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(57) Abstract: Compositions and methods for the therapy of malignant diseases, such as leukemia and cancer, are disclosed. The compositions comprise one or more of a WT1 polynucleotide, a WT1 polypeptide, an antigen-presenting cell presenting a WT1 polypeptide, an antibody that specifically binds to a WT1 polypeptide; or a T cell that specifically reacts with a WT1 polypeptide. Such compositions may be used, for example, for the prevention and treatment of metastatic diseases.

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COMPOSITIONS AND METHODS FOR WT1 SPECIFIC IMMUNOTHERAPY

TECHNICAL FIELD

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The present invention relates generally to the immunotherapy of malignant diseases such as leukemia and cancers. The invention is more specifically related to compositions for generating or enhancing an immune response to WT1, and to the use of such compositions for preventing and/or treating malignant diseases.

BACKGROUND OF THE INVENTION

Cancer and leukemia are significant health problems in the United States and throughout the world. Although advances have been made in detection and treatment of such diseases, no vaccine or other universally successful method for prevention or treatment of cancer and leukemia is currently available. Management of the diseases currently relies on a combination of early diagnosis and aggressive treatment, which may include one or more of a variety of treatments such as surgery, radiotherapy, chemotherapy and hormone therapy. The course of treatment for a particular cancer is often selected based on a variety of prognostic parameters, including an analysis of specific tumor markers. However, the use of established markers often leads to a result that is difficult to interpret, and the high mortality continues to be observed in many cancer patients.

Immunotherapies have the potential to substantially improve cancer and leukemia treatment and survival. Recent data demonstrate that leukemia can be cured by immunotherapy in the context of bone marrow transplantation (e.g., donor lymphocyte infusions). Such therapies may involve the generation or enhancement of an immune response to a tumor-associated antigen (TAA). However, to date, relatively few TAAs are known and the generation of an immune response against such antigens has, with rare exceptions, not been shown to be therapeutically beneficial.

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Accordingly, there is a need in the art for improved methods for leukemia and cancer prevention and therapy. The present invention fulfills these needs and further provides other related advantages.

SUMMARY OF THE INVENTION

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Briefly stated, this invention provides compositions and methods for the diagnosis and therapy of diseases such as leukemia and cancer. In one aspect, the present invention provides polypeptides comprising an immunogenic portion of a native WT1, or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera and/or T-cell lines or clones is not substantially diminished. Within certain embodiments, the polypeptide comprises no more than 16 consecutive amino acid residues of a native WT1 polypeptide. Within other embodiments, the polypeptide comprises an immunogenic portion of amino acid residues 1 - 174 of a native WT1 polypeptide or a variant thereof, wherein the polypeptide comprises no more than 16 consecutive amino acid residues present within amino acids 175 to 449 of the native WT1 polypeptide. The immunogenic portion preferably binds to an MHC class I and/or class II molecule. Within certain embodiments, the polypeptide comprises a sequence selected from the group consisting of (a) sequences recited in any one or more of Tables II - XLVI, (b) variants of the foregoing sequences that differ in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera and/or T-cell lines or clones is not substantially diminished and (c) mimetics of the polypeptides recited above, such that the ability of the mimetic to react with antigen-specific antisera and/or T cell lines or clones is not substantially diminished.

Within other embodiments, the polypeptide comprises a sequence selected from the group consisting of (a) ALLPAVPSL (SEQ ID NO:34), GATLKGVAA (SEQ ID NO:88), CMTWNQMNL (SEQ ID NOs: 49 and 258), SCLESQPTI (SEQ ID NOs: 199 and 296), SCLESQPAI (SEQ ID NO:198), NLYQMTSQL (SEQ ID NOs: 147 and 284),

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ALLPAVSSL (SEQ ID NOs: 35 and 255), RMFPNAPYL (SEQ ID NOs: 185 and 293), (b) variants of the foregoing sequences that differ in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera and/or T-cell lines or clones is not substantially diminished and (c) mimetics of the polypeptides recited above, such that the ability of the mimetic to react with antigen-specific antisera and/or T cell lines or clones is not substantially diminished. Mimetics may comprises amino acids in combination with one or more amino acid mimetics or may be entirely nonpeptide mimetics.

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Within further aspects, the present invention provides polypeptides comprising a variant of an immunogenic portion of a WT1 protein, wherein the variant differs from the immunogenic portion due to substitutions at between 1 and 3 amino acid positions within the immunogenic portion such that the ability of the variant to react with antigen-specific antisera and/or T-cell lines or clones is enhanced relative to a native WT1 protein.

The present invention further provides WT1 polynucleotides that encode a WT1 polypeptide as described above.

Within other aspects, the present invention provides pharmaceutical compositions and vaccines. Pharmaceutical compositions may comprise a polypeptide or mimetic as described above and/or one or more of (i) a WT1 polynucleotide; (ii) an antibody or antigen-binding fragment thereof that specifically binds to a WT1 polypeptide; (iii) a T cell that specifically reacts with a WT1 polypeptide or (iv) an antigen-presenting cell that expresses a WT1 polypeptide, in combination with a pharmaceutically acceptable carrier or excipient. Vaccines comprise a polypeptide as described above and/or one or more of (i) a WT1 polypucleotide, (ii) an antigen-presenting cell that expresses a WT1 polypeptide or (iii) an anti-idiotypic antibody, and a non-specific immune response enhancer. Within certain embodiments, less than 23 consecutive amino acid residues, preferably less than 17 amino acid residues, of a native WT1 polypeptide are present within a WT1 polypeptide employed within such pharmaceutical compositions and vaccines. The

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immune response enhancer may be an adjuvant. Preferably, an immune response enhancer enhances a T cell response.

The present invention further provides methods for enhancing or inducing an immune response in a patient, comprising administering to a patient a pharmaceutical composition or vaccine as described above. In certain embodiments, the patient is a human.

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The present invention further provides methods for inhibiting the development of a malignant disease in a patient, comprising administering to a patient a pharmaceutical composition or vaccine as described above. Malignant diseases include, but are not limited to leukemias (e.g., acute myeloid, acute lymphocytic and chronic myeloid) and cancers (e.g., breast, lung, thyroid or gastrointestinal cancer or a melanoma). The patient may, but need not, be afflicted with the malignant disease, and the administration of the pharmaceutical composition or vaccine may inhibit the onset of such a disease, or may inhibit progression and/or metastasis of an existing disease.

The present invention further provides, within other aspects, methods for removing cells expressing WT1 from bone marrow and/or peripheral blood or fractions thereof, comprising contacting bone marrow, peripheral blood or a fraction of bone marrow or peripheral blood with T cells that specifically react with a WT1 polypeptide, wherein the step of contacting is performed under conditions and for a time sufficient to permit the removal of WT1 positive cells to less than 10%, preferably less than 5% and more preferably less than 1%, of the number of myeloid or lymphatic cells in the bone marrow, peripheral blood or fraction. Bone marrow, peripheral blood and fractions may be obtained from a patient afflicted with a disease associated with WT1 expression, or may be obtained from a human or non-human mammal not afflicted with such a disease.

Within related aspects, the present invention provides methods for inhibiting the development of a malignant disease in a patient, comprising administering to a patient bone marrow, peripheral blood or a fraction of bone marrow or peripheral blood prepared as described above. Such bone marrow, peripheral blood or fractions may be autologous,

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or may be derived from a related or unrelated human or non-human animal (e.g., syngeneic or allogeneic).

In other aspects, the present invention provides methods for stimulating (or priming) and/or expanding T cells, comprising contacting T cells with a WT1 polypeptide under conditions and for a time sufficient to permit the stimulation and/or expansion of T cells. Such T cells may be autologous, allogeneic, syngeneic or unrelated WT1-specific T cells, and may be stimulated *in vitro* or *in vivo*. Expanded T cells may, within certain embodiments, be present within bone marrow, peripheral blood or a fraction of bone marrow or peripheral blood, and may (but need not) be clonal. Within certain embodiments, T cells may be present in a mammal during stimulation and/or expansion. WT1-specific T cells may be used, for example, within donor lymphocyte infusions.

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Within related aspects, methods are provided for inhibiting the development of a malignant disease in a patient, comprising administering to a patient T cells prepared as described above. Such T cells may, within certain embodiments, be autologous, syngeneic or allogeneic.

The present invention further provides, within other aspects, methods for monitoring the effectiveness of an immunization or therapy for a malignant disease associated with WT1 expression in a patient. Such methods are based on monitoring antibody, CD4+ T cell and/or CD8+ T cell responses in the patient. Within certain such aspects, a method may comprise the steps of: (a) incubating a first biological sample with one or more of: (i) a WT1 polypeptide; (ii) a polynucleotide encoding a WT1 polypeptide; or (iii) an antigen presenting cell that expresses a WT1 polypeptide, wherein the first biological sample is obtained from a patient prior to a therapy or immunization, and wherein the incubation is performed under conditions and for a time sufficient to allow immunocomplexes to form; (b) detecting immunocomplexes formed between the WT1 polypeptide and antibodies in the biological sample that specifically bind to the WT1 polypeptide; (c) repeating steps (a) and (b) using a second biological sample obtained from the same patient following therapy or immunization; and (d) comparing the number of

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immunocomplexes detected in the first and second biological samples, and therefrom monitoring the effectiveness of the therapy or immunization in the patient.

Within certain embodiments of the above methods, the step of detecting comprises (a) incubating the immunocomplexes with a detection reagent that is capable of binding to the immunocomplexes, wherein the detection reagent comprises a reporter group, (b) removing unbound detection reagent, and (c) detecting the presence or absence of the reporter group. The detection reagent may comprise, for example, a second antibody, or antigen-binding fragment thereof, capable of binding to the antibodies that specifically bind to the WT1 polypeptide or a molecule such as Protein A. Within other embodiments, a reporter group is bound to the WT1 polypeptide, and the step of detecting comprises removing unbound WT1 polypeptide and subsequently detecting the presence or absence of the reporter group.

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Within further aspects, methods for monitoring the effectiveness of an immunization or therapy for a malignant disease associated with WT1 expression in a patient may comprise the steps of: (a) incubating a first biological sample with one or more of: (i) a WT1 polypeptide; (ii) a polynucleotide encoding a WT1 polypeptide; or (iii) an antigen presenting cell that expresses a WT1 polypeptide, wherein the biological sample comprises CD4+ and/or CD8+ T cells and is obtained from a patient prior to a therapy or immunization, and wherein the incubation is performed under conditions and for a time sufficient to allow specific activation, proliferation and/or lysis of T cells; (b) detecting an amount of activation, proliferation and/or lysis of the T cells; (c) repeating steps (a) and (b) using a second biological sample comprising CD4+ and/or CD8+ T cells, wherein the second biological sample is obtained from the same patient following therapy or immunization; and (d) comparing the amount of activation, proliferation and/or lysis of T cells in the first and second biological samples, and therefrom monitoring the effectiveness of the therapy or immunization in the patient.

The present invention further provides methods for inhibiting the development of a malignant disease associated with WT1 expression in a patient,

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comprising the steps of: (a) incubating CD4⁺ and/or CD8+ T cells isolated from a patient with one or more of: (i) a WT1 polypeptide; (ii) a polynucleotide encoding a WT1 polypeptide; or (iii) an antigen presenting cell that expresses a WT1 polypeptide, such that the T cells proliferate; and (b) administering to the patient an effective amount of the proliferated T cells, and therefrom inhibiting the development of a malignant disease in the patient. Within certain embodiments, the step of incubating the T cells may be repeated one or more times.

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Within other aspects, the present invention provides methods for inhibiting the development of a malignant disease associated with WT1 expression in a patient, comprising the steps of: (a) incubating CD4⁺ and/or CD8+ T cells isolated from a patient with one or more of: (i) a WT1 polypeptide; (ii) a polynucleotide encoding a WT1 polypeptide; or (iii) an antigen presenting cell that expresses a WT1 polypeptide, such that the T cells proliferate; (b) cloning one or more cells that proliferated; and (c) administering to the patient an effective amount of the cloned T cells.

Within other aspects, methods are provided for determining the presence or absence of a malignant disease associated with WT1 expression in a patient, comprising the steps of: (a) incubating CD4⁺ and/or CD8+ T cells isolated from a patient with one or more of: (i) a WT1 polypeptide; (ii) a polynucleotide encoding a WT1 polypeptide; or (iii) an antigen presenting cell that expresses a WT1 polypeptide; and (b) detecting the presence or absence of specific activation of the T cells, therefrom determining the presence or absence of a malignant disease associated with WT1 expression. Within certain embodiments, the step of detecting comprises detecting the presence or absence of proliferation of the T cells.

Within further aspects, the present invention provides methods for determining the presence or absence of a malignant disease associated with WT1 expression in a patient, comprising the steps of: (a) incubating a biological sample obtained from a patient with one or more of: (i) a WT1 polypeptide; (ii) a polynucleotide encoding a WT1 polypeptide; or (iii) an antigen presenting cell that expresses a WT1

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polypeptide, wherein the incubation is performed under conditions and for a time sufficient to allow immunocomplexes to form; and (b) detecting immunocomplexes formed between the WT1 polypeptide and antibodies in the biological sample that specifically bind to the WT1 polypeptide; and therefrom determining the presence or absence of a malignant disease associated with WT1 expression.

These and other aspects of the present invention will become apparent upon reference to the following detailed description and attached drawings. All references disclosed herein are hereby incorporated by reference in their entirety as if each was incorporated individually.

10 BRIEF DESCRIPTION OF THE DRAWINGS

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Figure 1 depicts a comparison of the mouse (MO) and human (HU) WT1 protein sequences (SEQ ID NOS: 320 and 319 respectively).

Figure 2 is a Western blot illustrating the detection of WT1 specific antibodies in patients with hematological malignancy (AML). Lane 1 shows molecular weight markers; lane 2 shows a positive control (WT1 positive human leukemia cell line immunoprecipitated with a WT1 specific antibody); lane 3 shows a negative control (WT1 positive cell line immunoprecipitated with mouse sera); and lane 4 shows a WT1 positive cell line immunoprecipitated with sera of a patient with AML. For lanes 2-4, the immunoprecipitate was separated by gel electrophoresis and probed with a WT1 specific antibody.

Figure 3 is a Western blot illustrating the detection of a WT1 specific antibody response in B6 mice immunized with TRAMP-C, a WT1 positive tumor cell line. Lanes 1, 3 and 5 show molecular weight markers, and lanes 2, 4 and 6 show a WT1 specific positive control (N180, Santa Cruz Biotechnology, polypeptide spanning 180 amino acids of the N-terminal region of the WT1 protein, migrating on the Western blot at 52 kD). The primary antibody used was WT180 in lane 2, sera of non-immunized B6 mice in lane 4 and sera of the immunized B6 mice in lane 6.

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Figure 4 is a Western blot illustrating the detection of WT1 specific antibodies in mice immunized with representative WT1 peptides. Lanes 1, 3 and 5 show molecular weight markers and lanes 2, 4 and 6 show a WT1 specific positive control (N180, Santa Cruz Biotechnology, polypeptide spanning 180 amino acids of the N-terminal region of the WT1 protein, migrating on the Western blot at 52 kD). The primary antibody used was WT180 in lane 2, sera of non-immunized B6 mice in lane 4 and sera of the immunized B6 mice in lane 6.

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Figures 5A to 5C are graphs illustrating the stimulation of proliferative T cell responses in mice immunized with representative WT1 peptides. Thymidine incorporation assays were performed using one T cell line and two different clones, as indicated, and results were expressed as cpm. Controls indicated on the x axis were no antigen (No Ag) and B6/media; antigens used were p6-22 human (p1), p117-139 (p2) or p244-262 human (p3).

Figure 6A and 6B are histograms illustrating the stimulation of proliferative T cell responses in mice immunized with representative WT1 peptides. Three weeks after the third immunization, spleen cells of mice that had been inoculated with Vaccine A or Vaccine B were cultured with medium alone (medium) or spleen cells and medium (B6/no antigen), B6 spleen cells pulsed with the peptides p6-22 (p6), p117-139 (p117), p244-262 (p244) (Vaccine A; Figure 6A) or p287-301 (p287), p299-313 (p299), p421-435 (p421) (Vaccine B; Figure 6B) and spleen cells pulsed with an irrelevant control peptide (irrelevant peptide) at 25ug/ml and were assayed after 96hr for proliferation by (³H) thymidine incorporation. Bars represent the stimulation index (SI), which is calculated as the mean of the experimental wells divided by the mean of the control (B6 spleen cells with no antigen).

Figures 7A-7D are histograms illustrating the generation of proliferative T-cell lines and clones specific for p117-139 and p6-22. Following *in vivo* immunization, the initial three *in vitro* stimulations (IVS) were carried out using all three peptides of Vaccine A or B, respectively. Subsequent IVS were carried out as single peptide stimulations using

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only the two relevant peptides p117-139 and p6-22. Clones were derived from both the p6-22 and p117-139 specific T cell lines, as indicated. T cells were cultured with medium alone (medium) or spleen cells and medium (B6/no antigen), B6 spleen cells pulsed with the peptides p6-22 (p6), p117-139 (p117) or an irrelevant control peptide (irrelevant peptide) at 25ug/ml and were assayed after 96hr for proliferation by (³H) thymidine incorporation. Bars represent the stimulation index (SI), which is calculated as the mean of the experimental wells divided by the mean of the control (B6 spleen cells with no antigen).

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Figures 8A and 8B present the results of TSITES Analysis of human WT1 (SEQ ID NO:319) for peptides that have the potential to elicit Th responses. Regions indicated by "A" are AMPHI midpoints of blocks, "R" indicates residues matching the Rothbard/Taylor motif, "D" indicates residues matching the IAd motif, and 'd' indicates residues matching the IEd motif.

Figures 9A and 9B are graphs illustrating the elicitation of WT1 peptide-specific CTL in mice immunized with WT1 peptides. Figure 9A illustrates the lysis of target cells by allogeneic cell lines and Figure 9B shows the lysis of peptide coated cell lines. In each case, the % lysis (as determined by standard chromium release assays) is shown at three indicated effector:target ratios. Results are provided for lymphoma cells (LSTRA and E10), as well as E10 + p235-243 (E10+P235). E10 cells are also referred to herein as EL-4 cells.

Figures 10A-10D are graphs illustrating the elicitation of WT1 specific CTL, which kill WT1 positive tumor cell lines but do not kill WT1 negative cell lines, following vaccination of B6 mice with WT1 peptide P117. Figure 10A illustrates that T-cells of non-immunized B6 mice do not kill WT1 positive tumor cell lines. Figure 10B illustrates the lysis of the target cells by allogeneic cell lines. Figures 10C and 10D demonstrate the lysis of WT1 positive tumor cell lines, as compared to WT1 negative cell lines in two different experiments. In addition, Figures 10C and 10D show the lysis of peptide-coated cell lines (WT1 negative cell line E10 coated with the relevant WT1 peptide

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P117) In each case, the % lysis (as determined by standard chromium release assays) is shown at three indicated effector:target ratios. Results are provided for lymphoma cells (E10), prostate cancer cells (TRAMP-C), a transformed fibroblast cell line (BLK-SV40), as well as E10+p117.

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Figures 11A and 11B are histograms illustrating the ability of representative peptide P117-139 specific CTL to lyse WT1 positive tumor cells. Three weeks after the third immunization, spleen cells of mice that had been inoculated with the peptides p235-243 or p117-139 were stimulated in vitro with the relevant peptide and tested for ability to lyse targets incubated with WT1 peptides as well as WT1 positive and negative tumor cells. The bars represent the mean % specific lysis in chromium release assays performed in triplicate with an E:T ratio of 25:1. Figure 11A shows the cytotoxic activity of the p235-243 specific T cell line against the WT1 negative cell line EL-4 (EL-4, WT1 negative); EL-4 pulsed with the relevant (used for immunization as well as for restimulation) peptide p235-243 (EL-4+p235); EL-4 pulsed with the irrelevant peptides p117-139 (EL-4+p117), p126-134 (EL-4+p126) or p130-138 (EL-4+p130) and the WT1 positive tumor cells BLK-SV40 (BLK-SV40, WT1 positive) and TRAMP-C (TRAMP-C, WT1 positive), as indicated. Figure 11B shows cytotoxic activity of the p117-139 specific T cell line against EL-4; EL-4 pulsed with the relevant peptide P117-139 (EL-4+p117) and EL-4 pulsed with the irrelevant peptides p123-131 (EL-4+p123), or p128-136 (EL-4+p128); BLK-SV40 and TRAMP-C, as indicated.

Figures 12A and 12B are histograms illustrating the specificity of lysis of WT1 positive tumor cells, as demonstrated by cold target inhibition. The bars represent the mean % specific lysis in chromium release assays performed in triplicate with an E:T ratio of 25:1. Figure 12A shows the cytotoxic activity of the p117-139 specific T cell line against the WT1 negative cell line EL-4 (EL-4, WT1 negative); the WT1 positive tumor cell line TRAMP-C (TRAMP-C, WT1 positive); TRAMP-C cells incubated with a ten-fold excess (compared to the hot target) of EL-4 cells pulsed with the relevant peptide p117-139 (TRAMP-C + p117 cold target) without ⁵¹Cr labeling and TRAMP-C cells incubated with

EL-4 pulsed with an irrelevant peptide without ⁵¹Cr labeling (TRAMP-C + irrelevant cold target), as indicated. Figure 12B shows the cytotoxic activity of the p117-139 specific T cell line against the WT1 negative cell line EL-4 (EL-4, WT1 negative); the WT1 positive tumor cell line BLK-SV40 (BLK-SV40, WT1 positive); BLK-SV40 cells incubated with the relevant cold target (BLK-SV40 + p117 cold target) and BLK-SV40 cells incubated with the irrelevant cold target (BLK-SV40 + irrelevant cold target), as indicated.

Figures 13A-13C are histograms depicting an evaluation of the 9mer CTL epitope within p117-139. The p117-139 tumor specific CTL line was tested against peptides within aa117-139 containing or lacking an appropriate H-2^b class I binding motif and following restimulation with p126-134 or p130-138. The bars represent the mean % specific lysis in chromium release assays performed in triplicate with an E:T ratio of 25:1. Figure 13A shows the cytotoxic activity of the p117-139 specific T cell line against the WT1 negative cell line EL-4 (EL-4, WT1 negative) and EL-4 cells pulsed with the peptides p117-139 (EL-4 + p117), p119-127 (EL-4 + p119), p120-128 (EL-4 + p120), p123-131 (EL-4 + p123), p126-134 (EL-4 + p126), p128-136 (EL-4 + p128), and p130-138 (EL-4 + p130). Figure 13B shows the cytotoxic activity of the CTL line after restimulation with p126-134 against the WT1 negative cell line EL-4, EL-4 cells pulsed with p117-139 (EL-4 + p117), p126-134 (EL-4 + p126) and the WT1 positive tumor cell line TRAMP-C. Figure 13C shows the cytotoxic activity of the CTL line after restimulation with p130-138 against EL-4, EL-4 cells pulsed with p117-139 (EL-4 + p117), p130-138 (EL-4 + p130) and the WT1 positive tumor cell line TRAMP-C.

DETAILED DESCRIPTION OF THE INVENTION

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As noted above, the present invention is generally directed to compositions and methods for the immunotherapy and diagnosis of malignant diseases. The compositions described herein may include WT1 polypeptides, WT1 polynucleotides, antigen-presenting cells (APC, e.g., dendritic cells) that express a WT1 polypeptide, agents such as antibodies that bind to a WT1 polypeptide and/or immune system cells (e.g., T

cells) specific for WT1. WT1 Polypeptides of the present invention generally comprise at least a portion of a Wilms Tumor gene product (WT1) or a variant thereof. Nucleic acid sequences of the subject invention generally comprise a DNA or RNA sequence that encodes all or a portion of such a polypeptide, or that is complementary to such a sequence. Antibodies are generally immune system proteins, or antigen-binding fragments thereof, that are capable of binding to a portion of a WT1 polypeptide. T cells that may be employed within such compositions are generally T cells (e.g., CD4⁺ and/or CD8⁺) that are specific for a WT1 polypeptide. Certain methods described herein further employ antigen-presenting cells that express a WT1 polypeptide as provided herein.

The present invention is based on the discovery that an immune response raised against a Wilms Tumor (WT) gene product (e.g., WT1) can provide prophylactic and/or therapeutic benefit for patients afflicted with malignant diseases characterized by increased WT1 gene expression. Such diseases include, but are not limited to, leukemias (e.g., acute myeloid leukemia (AML), chronic myeloid leukemia (CML), acute lymphocytic leukemia (ALL) and childhood ALL), as well as many cancers such as lung, breast, thyroid and gastrointestinal cancers and melanomas. The WT1 gene was originally identified and isolated on the basis of a cytogenetic deletion at chromosome 11p13 in patients with Wilms' tumor (see Call et al., U.S. Patent No. 5,350,840). The gene consists of 10 exons and encodes a zinc finger transcription factor, and sequences of mouse and human WT1 proteins are provided in Figure 1 and SEQ ID NOs: 319 and 320.

WT1 POLYPEPTIDES

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Within the context of the present invention, a WT1 polypeptide is a polypeptide that comprises at least an immunogenic portion of a native WT1 (i.e., a WT1 protein expressed by an organism that is not genetically modified), or a variant thereof, as described herein. A WT1 polypeptide may be of any length, provided that it comprises at least an immunogenic portion of a native protein or a variant thereof. In other words, a WT1 polypeptide may be an oligopeptide (i.e., consisting of a relatively small number of

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amino acid residues, such as 8-10 residues, joined by peptide bonds), a full length WT1 protein (e.g., present within a human or non-human animal, such as a mouse) or a polypeptide of intermediate size. Within certain embodiments, the use of WT1 polypeptides that contain a small number of consecutive amino acid residues of a native WT1 polypeptide is preferred. Such polypeptides are preferred for certain uses in which the generation of a T cell response is desired. For example, such a WT1 polypeptide may contain less than 23, preferably no more than 18, and more preferably no more than 15 consecutive amino acid residues, of a native WT1 polypeptide. Polypeptides comprising nine consecutive amino acid residues of a native WT1 polypeptide are generally suitable Additional sequences derived from the native protein and/or for such purposes. heterologous sequences may be present within any WT1 polypeptide, and such sequences may (but need not) possess further immunogenic or antigenic properties. Polypeptides as provided herein may further be associated (covalently or noncovalently) with other polypeptide or non-polypeptide compounds.

An "immunogenic portion," as used herein is a portion of a polypeptide that is recognized (*i.e.*, specifically bound) by a B-cell and/or T-cell surface antigen receptor. Certain preferred immunogenic portions bind to an MHC class I or class II molecule. As used herein, an immunogenic portion is said to "bind to" an MHC class I or class II molecule if such binding is detectable using any assay known in the art. For example, the ability of a polypeptide to bind to MHC class I may be evaluated indirectly by monitoring the ability to promote incorporation of ¹²⁵I labeled β2-microglobulin (β2m) into MHC class I/β2m/peptide heterotrimeric complexes (*see* Parker et al., *J. Immunol. 152*:163, 1994). Alternatively, functional peptide competition assays that are known in the art may be employed. Certain immunogenic portions have one or more of the sequences recited within one or more of Tables II - XIV. Representative immunogenic portions include, but are not limited to, RDLNALLPAVPSLGGGG (human WT1 residues 6-22; SEQ ID NO:1), PSQASSGQARMFPNAPYLPSCLE (human and mouse WT1 residues 117-139; SEQ ID NOs: 2 and 3 respectively), GATLKGVAAGSSSSVKWTE (human WT1 residues 244-

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262; SEQ ID NO:4), GATLKGVAA (human WT1 residues 244-252; SEQ ID NO:88), CMTWNQMNL (human and mouse WT1 residues 235-243; SEQ ID NOs: 49 and 258 respectively), SCLESQPTI (mouse WT1 residues 136-144; SEO ID NO:296), SCLESQPAI (human WT1 residues 136-144; SEQ ID NO:198), NLYQMTSQL (human and mouse WT1 residues 225-233; SEQ ID NOs: 147 and 284 respectively); ALLPAVSSL (mouse WT1 residues 10-18; SEQ ID NO:255); or RMFPNAPYL (human and mouse WT1 residues 126-134; SEQ ID NOs: 185 and 293 respectively). Further immunogenic portions are provided herein, and others may generally be identified using well known techniques, such as those summarized in Paul, Fundamental Immunology, 3rd ed., 243-247 (Raven Press, 1993) and references cited therein. Representative techniques for identifying immunogenic portions include screening polypeptides for the ability to react with antigenspecific antisera and/or T-cell lines or clones. An immunogenic portion of a native WT1 polypeptide is a portion that reacts with such antisera and/or T-cells at a level that is not substantially less than the reactivity of the full length WT1 (e.g., in an ELISA and/or T-cell reactivity assay). In other words, an immunogenic portion may react within such assays at a level that is similar to or greater than the reactivity of the full length polypeptide. Such screens may generally be performed using methods well known to those of ordinary skill in the art, such as those described in Harlow and Lane, Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory, 1988.

Alternatively, immunogenic portions may be identified using computer analysis, such as the Tsites program (see Rothbard and Taylor, EMBO J. 7:93-100, 1988; Deavin et al., Mol. Immunol. 33:145-155, 1996), which searches for peptide motifs that have the potential to elicit Th responses. CTL peptides with motifs appropriate for binding to murine and human class I or class II MHC may be identified according to BIMAS (Parker et al., J. Immunol. 152:163, 1994) and other HLA peptide binding prediction analyses. To confirm immunogenicity, a peptide may be tested using an HLA A2 transgenic mouse model and/or an in vitro stimulation assay using dendritic cells, fibroblasts or peripheral blood cells.

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As noted above, a composition may comprise a variant of a native WT1 protein. A polypeptide "variant," as used herein, is a polypeptide that differs from a native polypeptide in one or more substitutions, deletions, additions and/or insertions, such that the immunogenicity of the polypeptide is retained (i.e., the ability of the variant to react with antigen-specific antisera and/or T-cell lines or clones is not substantially diminished relative to the native polypeptide). In other words, the ability of a variant to react with antigen-specific antisera and/or T-cell lines or clones may be enhanced or unchanged, relative to the native polypeptide, or may be diminished by less than 50%, and preferably less than 20%, relative to the native polypeptide. Such variants may generally be identified by modifying one of the above polypeptide sequences and evaluating the reactivity of the modified polypeptide with antisera and/or T-cells as described herein. It has been found, within the context of the present invention, that a relatively small number of substitutions (e.g., 1 to 3) within an immunogenic portion of a WT1 polypeptide may serve to enhance the ability of the polypeptide to elicit an immune response. Suitable substitutions may generally be identified by using computer programs, as described above, and the effect confirmed based on the reactivity of the modified polypeptide with antisera and/or T-cells Accordingly, within certain preferred embodiments, a WT1 as described herein. polypeptide comprises a variant in which 1 to 3 amino acid resides within an immunogenic portion are substituted such that the ability to react with antigen-specific antisera and/or Tcell lines or clones is statistically greater than that for the unmodified polypeptide. Such substitutions are preferably located within an MHC binding site of the polypeptide, which may be identified as described above. Preferred substitutions allow increased binding to MHC class I or class II molecules.

Certain variants contain conservative substitutions. A "conservative substitution" is one in which an amino acid is substituted for another amino acid that has similar properties, such that one skilled in the art of peptide chemistry would expect the secondary structure and hydropathic nature of the polypeptide to be substantially unchanged. Amino acid substitutions may generally be made on the basis of similarity in

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polarity, charge, solubility, hydrophobicity, hydrophilicity and/or the amphipathic nature of the residues. For example, negatively charged amino acids include aspartic acid and glutamic acid; positively charged amino acids include lysine and arginine; and amino acids with uncharged polar head groups having similar hydrophilicity values include leucine, isoleucine and valine; glycine and alanine; asparagine and glutamine; and serine, threonine, phenylalanine and tyrosine. Other groups of amino acids that may represent conservative changes include: (1) ala, pro, gly, glu, asp, gln, asn, ser, thr; (2) cys, ser, tyr, thr; (3) val, ile, leu, met, ala, phe; (4) lys, arg, his; and (5) phe, tyr, trp, his. A variant may also, or alternatively, contain nonconservative changes. Variants may also (or alternatively) be modified by, for example, the deletion or addition of amino acids that have minimal influence on the immunogenicity, secondary structure and hydropathic nature of the polypeptide.

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As noted above, WT1 polypeptides may be conjugated to a signal (or leader) sequence at the N-terminal end of the protein which co-translationally or post-translationally directs transfer of the protein. A polypeptide may also, or alternatively, be conjugated to a linker or other sequence for ease of synthesis, purification or identification of the polypeptide (e.g., poly-His), or to enhance binding of the polypeptide to a solid support. For example, a polypeptide may be conjugated to an immunoglobulin Fc region.

Within certain specific embodiments, a polypeptide may be a fusion protein that comprises multiple polypeptides as described herein, or that comprises at least one polypeptide as described herein and an unrelated sequence, such as a known tumor protein. A fusion partner may, for example, assist in providing T helper epitopes (an immunological fusion partner), preferably T helper epitopes recognized by humans, or may assist in expressing the protein (an expression enhancer) at higher yields than the native recombinant protein. Certain preferred fusion partners are both immunological and expression enhancing fusion partners. Other fusion partners may be selected so as to increase the solubility of the protein or to enable the protein to be targeted to desired

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intracellular compartments. Still further fusion partners include affinity tags, which facilitate purification of the protein.

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In certain embodiments, the present invention provides fusion proteins comprising a polypeptide disclosed herein together with at least one of the sequences disclosed in U.S. Patent Application No. 09/352,616, filed July 13, 1999, which is incorporated herein in its entirety. Preferably one or more of the following known prostate antigens may be employed in a fusion protein with WT1: prostate specific antigen (PSA); prostatic acid phosphatase (PAP); and prostate specific membrane antigen (PSMA). The protein sequences for PSMA, PAP and PSA are provided in SEQ ID NO: 327-329, respectively. In certain embodiments, the fusion proteins of the present invention comprise WT1, PSA, PAP and/or PSMA in combination with one or more of the following the inventive antigens: P501S (amino acid sequence provided in SEQ ID NO: 333); P703P (amino acid sequences provided in SEQ ID NO: 330-332); P704P (cDNA sequence provided in SEQ ID NO: 334); P712P (cDNA sequence provided in SEQ ID NO: 335); P775P (cDNA sequence provided in SEQ ID NO: 336); P776P (cDNA sequence provided in SEQ ID NO: 337); P790P (cDNA sequence provided in SEQ ID NO: 338). P711P (fulllength cDNA sequence provided in SEQ ID NO: 339, with the corresponding amino acid sequence provided in SEQ ID NO: 340); P710P (cDNA sequence provided in SEQ ID NO: 341-345); P714P (cDNA sequence provided in SEQ ID NO: 346); P510S (cDNA sequence provided in SEQ ID NO: 347); P774P (cDNA sequence provided in SEQ ID NO: 348); P705P (cDNA sequence provided in SEQ ID NO: 349, with the corresponding amino acid sequence provided in SEQ ID NO: 350); P503S (full-length cDNA sequence provided in SEQ ID NO: 351, with the corresponding amino acid sequence provided in SEQ ID NO: 352); P713P (cDNA sequence provided in SEQ ID NO: 353); P780P (cDNA sequence provided in SEQ ID NO: 354); P788P (cDNA sequence provided in SEQ ID NO: 355);

The cDNA sequence of a fusion protein for WT1 and PSA is provided in SEQ ID NO: 356 with the corresponding amino acid provided in SEQ ID NO: 357. In preferred embodiments, the inventive fusion proteins comprise one of the following

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combinations of antigens, or permutations: WT1 and PSA; WT1 and P703P; WT1 and P501S; WT1/P703P and P501S; WT1/PSA and P703P; WT1/PSA and P501S; WT1 and PAP; WT1/PAP and P703P; WT1/PAP and P501S; WT1 and PSMA; WT1/PSMA and P703P; WT1/PSMA and P501S; WT1/PSA/PAP and P501S; WT1/PSA/PAP and P501S; WT1/PSA/PAP and P703P; WT1/PSA/PSMA and P703P; WT1/PSA/PSMA and P501S; WT1/PSA/PSMA and P501S; WT1/PSA/PSMA and P501S; WT1/PSA/PAP/PSMA and P501S; WT1/PSA/PAP/PSMA and P501S. One of skill in the art will appreciate that the order of polypeptides within a fusion protein can be altered without substantially changing the therapeutic, prophylactic or diagnostic properties of the fusion protein.

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The fusion proteins described above are more immunogenic and will be effective in a greater number of prostate cancer patients than any of the individual components alone. The use of multiple antigens in the form of a fusion protein also lessens the likelihood of immunologic escape.

Fusion proteins may generally be prepared using standard techniques, including chemical conjugation. Preferably, a fusion protein is expressed as a recombinant protein, allowing the production of increased levels, relative to a non-fused protein, in an expression system. Briefly, DNA sequences encoding the polypeptide components may be assembled separately, and ligated into an appropriate expression vector. The 3' end of the DNA sequence encoding one polypeptide component is ligated, with or without a peptide linker, to the 5' end of a DNA sequence encoding the second polypeptide component so that the reading frames of the sequences are in phase. This permits translation into a single fusion protein that retains the biological activity of both component polypeptides.

A peptide linker sequence may be employed to separate the first and the second polypeptide components by a distance sufficient to ensure that each polypeptide folds into its secondary and tertiary structures. Such a peptide linker sequence is incorporated into the fusion protein using standard techniques well known in the art. Suitable peptide linker sequences may be chosen based on the following factors: (1) their ability to adopt a flexible extended conformation; (2) their inability to adopt a secondary

structure that could interact with functional epitopes on the first and second polypeptides; and (3) the lack of hydrophobic or charged residues that might react with the polypeptide functional epitopes. Preferred peptide linker sequences contain Gly, Asn and Ser residues. Other near neutral amino acids, such as Thr and Ala may also be used in the linker sequence. Amino acid sequences which may be usefully employed as linkers include those disclosed in Maratea et al., *Gene 40*:39-46, 1985; Murphy et al., *Proc. Natl. Acad. Sci. USA 83*:8258-8262, 1986; U.S. Patent No. 4,935,233 and U.S. Patent No. 4,751,180. The linker sequence may generally be from 1 to about 50 amino acids in length. Linker sequences are not required when the first and second polypeptides have non-essential N-terminal amino acid regions that can be used to separate the functional domains and prevent steric interference.

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The ligated DNA sequences are operably linked to suitable transcriptional or translational regulatory elements. The regulatory elements responsible for expression of DNA are located only 5' to the DNA sequence encoding the first polypeptides. Similarly, stop codons required to end translation and transcription termination signals are only present 3' to the DNA sequence encoding the second polypeptide.

Fusion proteins are also provided that comprise a polypeptide of the present invention together with an unrelated immunogenic protein. Preferably the immunogenic protein is capable of eliciting a recall response. Examples of such proteins include tetanus, tuberculosis and hepatitis proteins (see, for example, Stoute et al. New Engl. J. Med., 336:86-91, 1997).

Within preferred embodiments, an immunological fusion partner is derived from protein D, a surface protein of the gram-negative bacterium Haemophilus influenza B (WO 91/18926). Preferably, a protein D derivative comprises approximately the first third of the protein (e.g., the first N-terminal 100-110 amino acids), and a protein D derivative may be lipidated. Within certain preferred embodiments, the first 109 residues of a Lipoprotein D fusion partner is included on the N-terminus to provide the polypeptide with additional exogenous T-cell epitopes and to increase the expression level in E. coli (thus

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functioning as an expression enhancer). The lipid tail ensures optimal presentation of the antigen to antigen presenting cells. Other fusion partners include the non-structural protein from influenzae virus, NS1 (hemaglutinin). Typically, the N-terminal 81 amino acids are used, although different fragments that include T-helper epitopes may be used.

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In another embodiment, the immunological fusion partner is the protein known as LYTA, or a portion thereof (preferably a C-terminal portion). LYTA is derived from *Streptococcus pneumoniae*, which synthesizes an N-acetyl-L-alanine amidase known as amidase LYTA (encoded by the LytA gene; *Gene 43*:265-292, 1986). LYTA is an autolysin that specifically degrades certain bonds in the peptidoglycan backbone. The C-terminal domain of the LYTA protein is responsible for the affinity to the choline or to some choline analogues such as DEAE. This property has been exploited for the development of *E. coli* C-LYTA expressing plasmids useful for expression of fusion proteins. Purification of hybrid proteins containing the C-LYTA fragment at the amino terminus has been described (*see Biotechnology 10*:795-798, 1992). Within a preferred embodiment, a repeat portion of LYTA may be incorporated into a fusion protein. A repeat portion is found in the C-terminal region starting at residue 178. A particularly preferred repeat portion incorporates residues 188-305.

WT1 polypeptides may be prepared using any of a variety of well known techniques. Recombinant polypeptides encoded by a WT1 polynucleotide as described herein may be readily prepared from the polynucleotide. In general, any of a variety of expression vectors known to those of ordinary skill in the art may be employed to express recombinant WT1 polypeptides. Expression may be achieved in any appropriate host cell that has been transformed or transfected with an expression vector containing a DNA molecule that encodes a recombinant polypeptide. Suitable host cells include prokaryotes, yeast and higher eukaryotic cells. Preferably, the host cells employed are *E. coli*, yeast or a mammalian cell line such as COS or CHO. Supernatants from suitable host/vector systems which secrete recombinant protein or polypeptide into culture media may be first concentrated using a commercially available filter. The concentrate may then be applied to

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a suitable purification matrix such as an affinity matrix or an ion exchange resin. Finally, one or more reverse phase HPLC steps can be employed to further purify a recombinant polypeptide. Such techniques may be used to prepare native polypeptides or variants thereof. For example, polynucleotides that encode a variant of a native polypeptide may generally be prepared using standard mutagenesis techniques, such as oligonucleotide-directed site-specific mutagenesis, and sections of the DNA sequence may be removed to permit preparation of truncated polypeptides.

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Certain portions and other variants may also be generated by synthetic means, using techniques well known to those of ordinary skill in the art. For example, polypeptides having fewer than about 500 amino acids, preferably fewer than about 100 amino acids, and more preferably fewer than about 50 amino acids, may be synthesized. Polypeptides may be synthesized using any of the commercially available solid-phase techniques, such as the Merrifield solid-phase synthesis method, where amino acids are sequentially added to a growing amino acid chain. See Merrifield, J. Am. Chem. Soc. 85:2149-2146, 1963. Equipment for automated synthesis of polypeptides is commercially available from suppliers such as Applied BioSystems, Inc. (Foster City, CA), and may be operated according to the manufacturer's instructions.

In general, polypeptides and polynucleotides as described herein are isolated. An "isolated" polypeptide or polynucleotide is one that is removed from its original environment. For example, a naturally-occurring protein is isolated if it is separated from some or all of the coexisting materials in the natural system. Preferably, such polypeptides are at least about 90% pure, more preferably at least about 95% pure and most preferably at least about 99% pure. A polynucleotide is considered to be isolated if, for example, it is cloned into a vector that is not a part of the natural environment.

Within further aspects, the present invention provides mimetics of WT1 polypeptides. Such mimetics may comprise amino acids linked to one or more amino acid mimetics (i.e., one or more amino acids within the WT1 protein may be replaced by an amino acid mimetic) or may be entirely nonpeptide mimetics. An amino acid mimetic is a

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compound that is conformationally similar to an amino acid such that it can be substituted for an amino acid within a WT1 polypeptide without substantially diminishing the ability to react with antigen-specific antisera and/or T cell lines or clones. A nonpeptide mimetic is a compound that does not contain amino acids, and that has an overall conformation that is similar to a WT1 polypeptide such that the ability of the mimetic to react with WT1-specific antisera and/or T cell lines or clones is not substantially diminished relative to the ability of a WT1 polypeptide. Such mimetics may be designed based on standard techniques (e.g., nuclear magnetic resonance and computational techniques) that evaluate the three dimensional structure of a peptide sequence. Mimetics may be designed where one or more of the side chain functionalities of the WT1 polypeptide are replaced by groups that do not necessarily have the same size or volume, but have similar chemical and/or physical properties which produce similar biological responses. It should be understood that, within embodiments described herein, a mimetic may be substituted for a WT1 polypeptide.

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WT1 POLYNUCLEOTIDES

Any polynucleotide that encodes a WT1 polypeptide as described herein is a WT1 polynucleotide encompassed by the present invention. Such polynucleotides may be single-stranded (coding or antisense) or double-stranded, and may be DNA (genomic, cDNA or synthetic) or RNA molecules. Additional coding or non-coding sequences may, but need not, be present within a polynucleotide of the present invention, and a polynucleotide may, but need not, be linked to other molecules and/or support materials.

WT1 polynucleotides may encode a native WT1 protein, or may encode a variant of WT1 as described herein. Polynucleotide variants may contain one or more substitutions, additions, deletions and/or insertions such that the immunogenicity of the encoded polypeptide is not diminished, relative to a native WT1 protein. The effect on the immunogenicity of the encoded polypeptide may generally be assessed as described herein. Preferred variants contain nucleotide substitutions, deletions, insertions and/or additions at

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no more than 20%, preferably at no more than 10%, of the nucleotide positions that encode an immunogenic portion of a native WT1 sequence. Certain variants are substantially homologous to a native gene, or a portion thereof. Such polynucleotide variants are capable of hybridizing under moderately stringent conditions to a naturally occurring DNA sequence encoding a WT1 polypeptide (or a complementary sequence). Suitable moderately stringent conditions include prewashing in a solution of 5 X SSC, 0.5% SDS, 1.0 mM EDTA (pH 8.0); hybridizing at 50°C-65°C, 5 X SSC, overnight; followed by washing twice at 65°C for 20 minutes with each of 2X, 0.5X and 0.2X SSC containing 0.1% SDS). Such hybridizing DNA sequences are also within the scope of this invention.

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It will be appreciated by those of ordinary skill in the art that, as a result of the degeneracy of the genetic code, there are many nucleotide sequences that encode a WT1 polypeptide. Some of these polynucleotides bear minimal homology to the nucleotide sequence of any native gene. Nonetheless, polynucleotides that vary due to differences in codon usage are specifically contemplated by the present invention.

Once an immunogenic portion of WT1 is identified, as described above, a WT1 polynucleotide may be prepared using any of a variety of techniques. For example, a WT1 polynucleotide may be amplified from cDNA prepared from cells that express WT1. Such polynucleotides may be amplified via polymerase chain reaction (PCR). For this approach, sequence-specific primers may be designed based on the sequence of the immunogenic portion and may be purchased or synthesized. For example, suitable primers for PCR amplification of a human WT1 gene include: first step - P118: 1434-1414: 5' GAG AGT CAG ACT TGA AAG CAGT 3' (SEQ ID NO:5) and P135: 5' CTG AGC CTC AGC AAA TGG GC 3' (SEQ ID NO:6); second step - P136: 5' GAG CAT GCA TGG GCT CCG ACG TGC GGG 3' (SEQ ID NO:7) and P137: 5' GGG GTA CCC ACT GAA CGG TCC CCG A 3' (SEQ ID NO:8). Primers for PCR amplification of a mouse WT1 gene include: first step - P138: 5' TCC GAG CCG CAC CTC ATG 3' (SEQ ID NO:9) and P139: 5' GCC TGG GAT GCT GGA CTG 3' (SEQ ID NO:10), second step -

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P140: 5' GAG CAT GCG ATG GGT TCC GAC GTG CGG 3' (SEQ ID NO:11) and P141: 5' GGG GTA CCT CAA AGC GCC ACG TGG AGT TT 3' (SEQ ID NO:12).

An amplified portion may then be used to isolate a full length gene from a human genomic DNA library or from a suitable cDNA library, using well known techniques. Alternatively, a full length gene can be constructed from multiple PCR fragments. WT1 polynucleotides may also be prepared by synthesizing oligonucleotide components, and ligating components together to generate the complete polynucleotide.

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WT1 polynucleotides may also be synthesized by any method known in the art, including chemical synthesis (e.g., solid phase phosphoramidite chemical synthesis). Modifications in a polynucleotide sequence may also be introduced using standard mutagenesis techniques, such as oligonucleotide-directed site-specific mutagenesis (see Adelman et al., DNA 2:183, 1983). Alternatively, RNA molecules may be generated by in vitro or in vivo transcription of DNA sequences encoding a WT1 polypeptide, provided that the DNA is incorporated into a vector with a suitable RNA polymerase promoter (such as T7 or SP6). Certain portions may be used to prepare an encoded polypeptide, as described herein. In addition, or alternatively, a portion may be administered to a patient such that the encoded polypeptide is generated in vivo (e.g., by transfecting antigen-presenting cells such as dendritic cells with a cDNA construct encoding a WT1 polypeptide, and administering the transfected cells to the patient).

Polynucleotides that encode a WT1 polypeptide may generally be used for production of the polypeptide, *in vitro* or *in vivo*. WT1 polynucleotides that are complementary to a coding sequence (*i.e.*, antisense polynucleotides) may also be used as a probe or to inhibit WT1 expression. cDNA constructs that can be transcribed into antisense RNA may also be introduced into cells of tissues to facilitate the production of antisense RNA.

Any polynucleotide may be further modified to increase stability *in vivo*. Possible modifications include, but are not limited to, the addition of flanking sequences at the 5' and/or 3' ends; the use of phosphorothioate or 2' O-methyl rather than

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phosphodiesterase linkages in the backbone; and/or the inclusion of nontraditional bases such as inosine, queosine and wybutosine, as well as acetyl- methyl-, thio- and other modified forms of adenine, cytidine, guanine, thymine and uridine.

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Nucleotide sequences as described herein may be joined to a variety of other nucleotide sequences using established recombinant DNA techniques. For example, a polynucleotide may be cloned into any of a variety of cloning vectors, including plasmids, phagemids, lambda phage derivatives and cosmids. Vectors of particular interest include expression vectors, replication vectors, probe generation vectors and sequencing vectors. In general, a vector will contain an origin of replication functional in at least one organism, convenient restriction endonuclease sites and one or more selectable markers. Other elements will depend upon the desired use, and will be apparent to those of ordinary skill in the art.

Within certain embodiments, polynucleotides may be formulated so as to permit entry into a cell of a mammal, and expression therein. Such formulations are particularly useful for therapeutic purposes, as described below. Those of ordinary skill in the art will appreciate that there are many ways to achieve expression of a polynucleotide in a target cell, and any suitable method may be employed. For example, a polynucleotide may be incorporated into a viral vector such as, but not limited to, adenovirus, adeno-associated virus, retrovirus, or vaccinia or other pox virus (e.g., avian pox virus). Techniques for incorporating DNA into such vectors are well known to those of ordinary skill in the art. A retroviral vector may additionally transfer or incorporate a gene for a selectable marker (to aid in the identification or selection of transduced cells) and/or a targeting moiety, such as a gene that encodes a ligand for a receptor on a specific target cell, to render the vector target specific. Targeting may also be accomplished using an antibody, by methods known to those of ordinary skill in the art. cDNA constructs within such a vector may be used, for example, to transfect human or animal cell lines for use in establishing WT1 positive tumor models which may be used to perform tumor protection

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and adoptive immunotherapy experiments to demonstrate tumor or leukemia-growth inhibition or lysis of such cells.

Other therapeutic formulations for polynucleotides include colloidal dispersion systems, such as macromolecule complexes, nanocapsules, microspheres, beads, and lipid-based systems including oil-in-water emulsions, micelles, mixed micelles, and liposomes. A preferred colloidal system for use as a delivery vehicle *in vitro* and *in vivo* is a liposome (*i.e.*, an artificial membrane vesicle). The preparation and use of such systems is well known in the art.

10 ANTIBODIES AND FRAGMENTS THEREOF

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The present invention further provides binding agents, such as antibodies and antigen-binding fragments thereof, that specifically bind to a WT1 polypeptide. As used herein, an agent is said to "specifically bind" to a WT1 polypeptide if it reacts at a detectable level (within, for example, an ELISA) with a WT1 polypeptide, and does not react detectably with unrelated proteins under similar conditions. As used herein, "binding" refers to a noncovalent association between two separate molecules such that a "complex" is formed. The ability to bind may be evaluated by, for example, determining a binding constant for the formation of the complex. The binding constant is the value obtained when the concentration of the complex is divided by the product of the component concentrations. In general, two compounds are said to "bind," in the context of the present invention, when the binding constant for complex formation exceeds about 10³ L/mol. The binding constant maybe determined using methods well known in the art.

Any agent that satisfies the above requirements may be a binding agent. In a preferred embodiment, a binding agent is an antibody or an antigen-binding fragment thereof. Certain antibodies are commercially available from, for example, Santa Cruz Biotechnology (Santa Cruz, CA). Alternatively, antibodies may be prepared by any of a variety of techniques known to those of ordinary skill in the art. See, e.g., Harlow and Lane, Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory, 1988. In

general, antibodies can be produced by cell culture techniques, including the generation of monoclonal antibodies as described herein, or via transfection of antibody genes into suitable bacterial or mammalian cell hosts, in order to allow for the production of recombinant antibodies. In one technique, an immunogen comprising the polypeptide is initially injected into any of a wide variety of mammals (e.g., mice, rats, rabbits, sheep or goats). In this step, the polypeptides of this invention may serve as the immunogen without modification. Alternatively, particularly for relatively short polypeptides, a superior immune response may be elicited if the polypeptide is joined to a carrier protein, such as bovine serum albumin or keyhole limpet hemocyanin. The immunogen is injected into the animal host, preferably according to a predetermined schedule incorporating one or more booster immunizations, and the animals are bled periodically. Polyclonal antibodies specific for the polypeptide may then be purified from such antisera by, for example, affinity chromatography using the polypeptide coupled to a suitable solid support.

Monoclonal antibodies specific for the antigenic polypeptide of interest may be prepared, for example, using the technique of Kohler and Milstein, Eur. J. Immunol. 6:511-519, 1976, and improvements thereto. Briefly, these methods involve the preparation of immortal cell lines capable of producing antibodies having the desired specificity (i.e., reactivity with the polypeptide of interest). Such cell lines may be produced, for example, from spleen cells obtained from an animal immunized as described above. The spleen cells are then immortalized by, for example, fusion with a myeloma cell fusion partner, preferably one that is syngeneic with the immunized animal. A variety of fusion techniques may be employed. For example, the spleen cells and myeloma cells may be combined with a nonionic detergent for a few minutes and then plated at low density on a selective medium that supports the growth of hybrid cells, but not myeloma cells. A preferred selection technique uses HAT (hypoxanthine, aminopterin, thymidine) selection. After a sufficient time, usually about 1 to 2 weeks, colonies of hybrids are observed. Single colonies are selected and their culture supernatants tested for binding activity against the polypeptide. Hybridomas having high reactivity and specificity are preferred.

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Monoclonal antibodies may be isolated from the supernatants of growing hybridoma colonies. In addition, various techniques may be employed to enhance the yield, such as injection of the hybridoma cell line into the peritoneal cavity of a suitable vertebrate host, such as a mouse. Monoclonal antibodies may then be harvested from the ascites fluid or the blood. Contaminants may be removed from the antibodies by conventional techniques, such as chromatography, gel filtration, precipitation, and extraction. The polypeptides of this invention may be used in the purification process in, for example, an affinity chromatography step.

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Within certain embodiments, the use of antigen-binding fragments of antibodies may be preferred. Such fragments include Fab fragments, which may be prepared using standard techniques. Briefly, immunoglobulins may be purified from rabbit serum by affinity chromatography on Protein A bead columns (Harlow and Lane, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, 1988) and digested by papain to yield Fab and Fc fragments. The Fab and Fc fragments may be separated by affinity chromatography on protein A bead columns.

Monoclonal antibodies and fragments thereof may be coupled to one or more therapeutic agents. Suitable agents in this regard include radioactive tracers and chemotherapeutic agents, which may be used, for example, to purge autologous bone marrow *in vitro*). Representative therapeutic agents include radionuclides, differentiation inducers, drugs, toxins, and derivatives thereof. Preferred radionuclides include ⁹⁰Y, ¹²³I, ¹²⁵I, ¹³¹I, ¹⁸⁶Re, ¹⁸⁸Re, ²¹¹At, and ²¹²Bi. Preferred drugs include methotrexate, and pyrimidine and purine analogs. Preferred differentiation inducers include phorbol esters and butyric acid. Preferred toxins include ricin, abrin, diptheria toxin, cholera toxin, gelonin, Pseudomonas exotoxin, Shigella toxin, and pokeweed antiviral protein. For diagnostic purposes, coupling of radioactive agents may be used to facilitate tracing of metastases or to determine the location of WT1-positive tumors.

A therapeutic agent may be coupled (e.g., covalently bonded) to a suitable monoclonal antibody either directly or indirectly (e.g., via a linker group). A direct

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reaction between an agent and an antibody is possible when each possesses a substituent capable of reacting with the other. For example, a nucleophilic group, such as an amino or sulfhydryl group, on one may be capable of reacting with a carbonyl-containing group, such as an anhydride or an acid halide, or with an alkyl group containing a good leaving group (e.g., a halide) on the other.

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Alternatively, it may be desirable to couple a therapeutic agent and an antibody via a linker group. A linker group can function as a spacer to distance an antibody from an agent in order to avoid interference with binding capabilities. A linker group can also serve to increase the chemical reactivity of a substituent on an agent or an antibody, and thus increase the coupling efficiency. An increase in chemical reactivity may also facilitate the use of agents, or functional groups on agents, which otherwise would not be possible.

It will be evident to those skilled in the art that a variety of bifunctional or polyfunctional reagents, both homo- and hetero-functional (such as those described in the catalog of the Pierce Chemical Co., Rockford, IL), may be employed as the linker group. Coupling may be effected, for example, through amino groups, carboxyl groups, sulfhydryl groups or oxidized carbohydrate residues. There are numerous references describing such methodology, e.g., U.S. Patent No. 4,671,958, to Rodwell et al.

Where a therapeutic agent is more potent when free from the antibody portion of the immunoconjugates of the present invention, it may be desirable to use a linker group which is cleavable during or upon internalization into a cell. A number of different cleavable linker groups have been described. The mechanisms for the intracellular release of an agent from these linker groups include cleavage by reduction of a disulfide bond (e.g., U.S. Patent No. 4,489,710, to Spitler), by irradiation of a photolabile bond (e.g., U.S. Patent No. 4,625,014, to Senter et al.), by hydrolysis of derivatized amino acid side chains (e.g., U.S. Patent No. 4,638,045, to Kohn et al.), by serum complement-mediated hydrolysis (e.g., U.S. Patent No. 4,671,958, to Rodwell et al.), and acid-catalyzed hydrolysis (e.g., U.S. Patent No. 4,569,789, to Blattler et al.).

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It may be desirable to couple more than one agent to an antibody. In one embodiment, multiple molecules of an agent are coupled to one antibody molecule. In another embodiment, more than one type of agent may be coupled to one antibody. Regardless of the particular embodiment, immunoconjugates with more than one agent may be prepared in a variety of ways. For example, more than one agent may be coupled directly to an antibody molecule, or linkers which provide multiple sites for attachment can be used. Alternatively, a carrier can be used. A carrier may bear the agents in a variety of ways, including covalent bonding either directly or via a linker group. Suitable carriers include proteins such as albumins (e.g., U.S. Patent No. 4,507,234, to Kato et al.), peptides and polysaccharides such as aminodextran (e.g., U.S. Patent No. 4,699,784, to Shih et al.). A carrier may also bear an agent by noncovalent bonding or by encapsulation, such as within a liposome vesicle (e.g., U.S. Patent Nos. 4,429,008 and 4,873,088). Carriers specific for radionuclide agents include radiohalogenated small molecules and chelating compounds. For example, U.S. Patent No. 4,735,792 discloses representative radiohalogenated small molecules and their synthesis. A radionuclide chelate may be formed from chelating compounds that include those containing nitrogen and sulfur atoms as the donor atoms for binding the metal, or metal oxide, radionuclide. For example, U.S. Patent No. 4,673,562, to Davison et al. discloses representative chelating compounds and their synthesis.

A variety of routes of administration for the antibodies and immunoconjugates may be used. Typically, administration will be intravenous, intramuscular, subcutaneous or in the bed of a resected tumor. It will be evident that the precise dose of the antibody/immunoconjugate will vary depending upon the antibody used, the antigen density on the tumor, and the rate of clearance of the antibody.

Also provided herein are anti-idiotypic antibodies that mimic an immunogenic portion of WT1. Such antibodies may be raised against an antibody, or antigen-binding fragment thereof, that specifically binds to an immunogenic portion of WT1, using well known techniques. Anti-idiotypic antibodies that mimic an immunogenic

portion of WT1 are those antibodies that bind to an antibody, or antigen-binding fragment thereof, that specifically binds to an immunogenic portion of WT1, as described herein.

T CELLS

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Immunotherapeutic compositions may also, or alternatively, comprise T cells specific for WT1. Such cells may generally be prepared *in vitro* or *ex vivo*, using standard procedures. For example, T cells may be present within (or isolated from) bone marrow, peripheral blood or a fraction of bone marrow or peripheral blood of a mammal, such as a patient, using a commercially available cell separation system, such as the CEPRATETM system, available from CellPro Inc., Bothell WA (see also U.S. Patent No. 5,240,856; U.S. Patent No. 5,215,926; WO 89/06280; WO 91/16116 and WO 92/07243). Alternatively, T cells may be derived from related or unrelated humans, non-human animals, cell lines or cultures.

T cells may be stimulated with WT1 polypeptide, polynucleotide encoding a WT1 polypeptide and/or an antigen presenting cell (APC) that expresses a WT1 polypeptide. Such stimulation is performed under conditions and for a time sufficient to permit the generation of T cells that are specific for the WT1 polypeptide. Preferably, a WT1 polypeptide or polynucleotide is present within a delivery vehicle, such as a microsphere, to facilitate the generation of antigen-specific T cells. Briefly, T cells, which may be isolated from a patient or a related or unrelated donor by routine techniques (such as by Ficoll/Hypaque density gradient centrifugation of peripheral blood lymphocytes), are incubated with WT1 polypeptide. For example, T cells may be incubated *in vitro* for 2-9 days (typically 4 days) at 37°C with WT1 polypeptide (e.g., 5 to 25 μg/ml) or cells synthesizing a comparable amount of WT1 polypeptide. It may be desirable to incubate a separate aliquot of a T cell sample in the absence of WT1 polypeptide to serve as a control.

T cells are considered to be specific for a WT1 polypeptide if the T cells kill target cells coated with a WT1 polypeptide or expressing a gene encoding such a polypeptide. T cell specificity may be evaluated using any of a variety of standard

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techniques. For example, within a chromium release assay or proliferation assay, a stimulation index of more than two fold increase in lysis and/or proliferation, compared to negative controls, indicates T cell specificity. Such assays may be performed, for example, as described in Chen et al., Cancer Res. 54:1065-1070, 1994. Alternatively, detection of the proliferation of T cells may be accomplished by a variety of known techniques. For example, T cell proliferation can be detected by measuring an increased rate of DNA synthesis (e.g., by pulse-labeling cultures of T cells with tritiated thymidine and measuring the amount of tritiated thymidine incorporated into DNA). Other ways to detect T cell proliferation include measuring increases in interleukin-2 (IL-2) production, Ca²⁺ flux, or dye uptake, such as 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl-tetrazolium. Alternatively, synthesis of lymphokines (such as interferon-gamma) can be measured or the relative number of T cells that can respond to a WT1 polypeptide may be quantified. Contact with a WT1 polypeptide (200 ng/ml - 100 μg/ml, preferably 100 ng/ml - 25 μg/ml) for 3 - 7 days should result in at least a two fold increase in proliferation of the T cells and/or contact as described above for 2-3 hours should result in activation of the T cells, as measured using standard cytokine assays in which a two fold increase in the level of cytokine release (e.g., TNF or IFN-y) is indicative of T cell activation (see Coligan et al., Current Protocols in Immunology, vol. 1, Wiley Interscience (Greene 1998). WT1 specific T cells may be expanded using standard techniques. Within preferred embodiments, the T cells are derived from a patient or a related or unrelated donor and are administered to the patient following stimulation and expansion.

T cells that have been activated in response to a WT1 polypeptide, polynucleotide or WT1-expressing APC may be CD4⁺ and/or CD8⁺. Specific activation of CD4⁺ or CD8⁺ T cells may be detected in a variety of ways. Methods for detecting specific T cell activation include detecting the proliferation of T cells, the production of cytokines (e.g., lymphokines), or the generation of cytolytic activity (i.e., generation of cytotoxic T cells specific for WT1). For CD4⁺ T cells, a preferred method for detecting specific T cell activation is the detection of the proliferation of T cells. For CD8⁺ T cells, a

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preferred method for detecting specific T cell activation is the detection of the generation of cytolytic activity.

For therapeutic purposes, CD4⁺ or CD8⁺ T cells that proliferate in response to the WT1 polypeptide, polynucleotide or APC can be expanded in number either in vitro or in vivo. Proliferation of such T cells in vitro may be accomplished in a variety of ways. For example, the T cells can be re-exposed to WT1 polypeptide, with or without the addition of T cell growth factors, such as interleukin-2, and/or stimulator cells that synthesize a WT1 polypeptide. The addition of stimulator cells is preferred where generating CD8+ T cell responses. T cells can be grown to large numbers in vitro with retention of specificity in response to intermittent restimulation with WT1 polypeptide. Briefly, for the primary in vitro stimulation (IVS), large numbers of lymphocytes (e.g., greater than 4 x 10⁷) may be placed in flasks with media containing human serum. WT1 polypeptide (e.g., peptide at 10 µg/ml) may be added directly, along with tetanus toxoid (e.g., 5 µg/ml). The flasks may then be incubated (e.g., 37°C for 7 days). For a second IVS, T cells are then harvested and placed in new flasks with $2-3 \times 10^7$ irradiated peripheral blood mononuclear cells. WT1 polypeptide (e.g., 10 µg/ml) is added directly. The flasks are incubated at 37°C for 7 days. On day 2 and day 4 after the second IVS, 2-5 units of interleukin-2 (IL-2) may be added. For a third IVS, the T cells may be placed in wells and stimulated with the individual's own EBV transformed B cells coated with the peptide. IL-2 may be added on days 2 and 4 of each cycle. As soon as the cells are shown to be specific cytotoxic T cells, they may be expanded using a 10 day stimulation cycle with higher IL-2 (20 units) on days 2, 4 and 6.

Alternatively, one or more T cells that proliferate in the presence of WT1 polypeptide can be expanded in number by cloning. Methods for cloning cells are well known in the art, and include limiting dilution. Responder T cells may be purified from the peripheral blood of sensitized patients by density gradient centrifugation and sheep red cell rosetting and established in culture by stimulating with the nominal antigen in the presence of irradiated autologous filler cells. In order to generate CD4⁺ T cell lines, WT1

polypeptide is used as the antigenic stimulus and autologous peripheral blood lymphocytes (PBL) or lymphoblastoid cell lines (LCL) immortalized by infection with Epstein Barr virus are used as antigen presenting cells. In order to generate CD8⁺ T cell lines, autologous antigen-presenting cells transfected with an expression vector which produces WT1 polypeptide may be used as stimulator cells. Established T cell lines may be cloned 2-4 days following antigen stimulation by plating stimulated T cells at a frequency of 0.5 cells per well in 96-well flat-bottom plates with 1 x 10⁶ irradiated PBL or LCL cells and recombinant interleukin-2 (rIL2) (50 U/ml). Wells with established clonal growth may be identified at approximately 2-3 weeks after initial plating and restimulated with appropriate antigen in the presence of autologous antigen-presenting cells, then subsequently expanded by the addition of low doses of rIL2 (10 U/ml) 2-3 days following antigen stimulation. T cell clones may be maintained in 24-well plates by periodic restimulation with antigen and rIL2 approximately every two weeks.

Within certain embodiments, allogeneic T-cells may be primed (i.e., sensitized to WT1) in vivo and/or in vitro. Such priming may be achieved by contacting T cells with a WT1 polypeptide, a polynucleotide encoding such a polypeptide or a cell producing such a polypeptide under conditions and for a time sufficient to permit the priming of T cells. In general, T cells are considered to be primed if, for example, contact with a WT1 polypeptide results in proliferation and/or activation of the T cells, as measured by standard proliferation, chromium release and/or cytokine release assays as described herein. A stimulation index of more than two fold increase in proliferation or lysis, and more than three fold increase in the level of cytokine, compared to negative controls, indicates T-cell specificity. Cells primed in vitro may be employed, for example, within a bone marrow transplantation or as donor lymphocyte infusion.

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PHARMACEUTICAL COMPOSITIONS AND VACCINES

Within certain aspects, polypeptides, polynucleotides, antibodies and/or T cells may be incorporated into pharmaceutical compositions or vaccines. Alternatively, a

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pharmaceutical composition may comprise an antigen-presenting cell (e.g., a dendritic cell) transfected with a WT1 polynucleotide such that the antigen presenting cell expresses a WT1 polypeptide. Pharmaceutical compositions comprise one or more such compounds or cells and a physiologically acceptable carrier or excipient. Certain vaccines may comprise one or more such compounds or cells and a non-specific immune response enhancer, such as an adjuvant or a liposome (into which the compound is incorporated). Pharmaceutical compositions and vaccines may additionally contain a delivery system, such as biodegradable microspheres which are disclosed, for example, in U.S. Patent Nos. 4,897,268 and 5,075,109. Pharmaceutical compositions and vaccines within the scope of the present invention may also contain other compounds, which may be biologically active or inactive.

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Within certain embodiments, pharmaceutical compositions and vaccines are designed to elicit T cell responses specific for a WT1 polypeptide in a patient, such as a human. In general, T cell responses may be favored through the use of relatively short polypeptides (e.g., comprising less than 23 consecutive amino acid residues of a native WT1 polypeptide, preferably 4-16 consecutive residues, more preferably 8-16 consecutive residues and still more preferably 8-10 consecutive residues. Alternatively, or in addition, a vaccine may comprise a non-specific immune response enhancer that preferentially enhances a T cell response. In other words, the immune response enhancer may enhance the level of a T cell response to a WT1 polypeptide by an amount that is proportionally greater than the amount by which an antibody response is enhanced. For example, when compared to a standard oil based adjuvant, such as CFA, an immune response enhancer that preferentially enhances a T cell response may enhance a proliferative T cell response by at least two fold, a lytic response by at least 10%, and/or T cell activation by at least two fold compared to WT1-megative control cell lines, while not detectably enhancing an antibody response. The amount by which a T cell or antibody response to a WT1 polypeptide is enhanced may generally be determined using any representative technique known in the art, such as the techniques provided herein.

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A pharmaceutical composition or vaccine may contain DNA encoding one or more of the polypeptides as described above, such that the polypeptide is generated in situ. As noted above, the DNA may be present within any of a variety of delivery systems known to those of ordinary skill in the art, including nucleic acid expression systems, bacterial and viral expression systems and mammalian expression systems. Appropriate nucleic acid expression systems contain the necessary DNA, cDNA or RNA sequences for expression in the patient (such as a suitable promoter and terminating signal). Bacterial delivery systems involve the administration of a bacterium (such as Bacillus-Calmette-Guerrin) that expresses an immunogenic portion of the polypeptide on its cell surface. In a preferred embodiment, the DNA may be introduced using a viral expression system (e.g., vaccinia or other pox virus, retrovirus, or adenovirus), which may involve the use of a nonpathogenic (defective), replication competent virus. Techniques for incorporating DNA into such expression systems are well known to those of ordinary skill in the art. The DNA may also be "naked," as described, for example, in Ulmer et al., Science 259:1745-1749, 1993 and reviewed by Cohen, Science 259:1691-1692, 1993. The uptake of naked DNA may be increased by coating the DNA onto biodegradable beads, which are efficiently transported into the cells.

As noted above, a pharmaceutical composition or vaccine may comprise an antigen-presenting cell that expresses a WT1 polypeptide. For therapeutic purposes, as described herein, the antigen presenting cell is preferably an autologous dendritic cell. Such cells may be prepared and transfected using standard techniques, such as those described by Reeves et al., *Cancer Res.* 56:5672-5677, 1996; Tuting et al., *J. Immunol.* 160:1139-1147, 1998; and Nair et al., *Nature Biotechnol.* 16:364-369, 1998). Expression of a WT1 polypeptide on the surface of an antigen-presenting cell may be confirmed by *in vitro* stimulation and standard proliferation as well as chromium release assays, as described herein.

While any suitable carrier known to those of ordinary skill in the art may be employed in the pharmaceutical compositions of this invention, the type of carrier will vary

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depending on the mode of administration. Compositions of the present invention may be formulated for any appropriate manner of administration, including for example, topical, oral, nasal, intravenous, intracranial, intraperitoneal, subcutaneous or intramuscular administration. For parenteral administration, such as subcutaneous injection, the carrier preferably comprises water, saline, alcohol, a fat, a wax or a buffer. For oral administration, any of the above carriers or a solid carrier, such as mannitol, lactose, starch, magnesium stearate, sodium saccharine, talcum, cellulose, glucose, sucrose, and magnesium carbonate, may be employed. Biodegradable microspheres (e.g., polylactate polyglycolate) may also be employed as carriers for the pharmaceutical compositions of this invention. For certain topical applications, formulation as a cream or lotion, using well known components, is preferred.

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Such compositions may also comprise buffers (e.g., neutral buffered saline or phosphate buffered saline), carbohydrates (e.g., glucose, mannose, sucrose or dextrans), mannitol, proteins, polypeptides or amino acids such as glycine, antioxidants, chelating agents such as EDTA or glutathione, adjuvants (e.g., aluminum hydroxide) and/or preservatives. Alternatively, compositions of the present invention may be formulated as a lyophilizate. Compounds may also be encapsulated within liposomes using well known technology.

Any of a variety of non-specific immune response enhancers, such as adjuvants, may be employed in the vaccines of this invention. Most adjuvants contain a substance designed to protect the antigen from rapid catabolism, such as aluminum hydroxide or mineral oil, and a stimulator of immune responses, such as lipid A, *Bortadella pertussis* or *Mycobacterium tuberculosis* derived proteins. Suitable non-specific immune response enhancers include alum-based adjuvants (e.g., Alhydrogel, Rehydragel, aluminum phosphate, Algammulin, aluminum hydroxide); oil based adjuvants (Freund's adjuvant (FA), Specol, RIBI, TiterMax, Montanide ISA50 or Seppic MONTANIDE ISA 720; cytokines (e.g., GM-CSF or Flat3-ligand); microspheres; nonionic block copolymer-based adjuvants; dimethyl dioctadecyl ammoniumbromide (DDA) based adjuvants AS-1, AS-2

(Smith Kline Beecham); Ribi Adjuvant system based adjuvants; QS21 (Aquila); saponin based adjuvants (crude saponin, the saponin Quil A); muramyl dipeptide (MDP) based adjuvants such as SAF (Syntex adjuvant in its microfluidized form (SAF-m)); dimethyl-dioctadecyl ammonium bromide (DDA); human complement based adjuvants *m. vaccae* and derivatives; immune stimulating complex (iscom) based adjuvants; inactivated toxins; and attenuated infectious agents (such as *M. tuberculosis*).

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As noted above, within certain embodiments, immune response enhancers are chosen for their ability to preferentially elicit or enhance a T cell response (e.g., CD4⁺ and/or CD8⁺) to a WT1 polypeptide. Such immune response enhancers are well known in the art, and include (but are not limited to) Montanide ISA50, Seppic MONTANIDE ISA 720, cytokines (e.g., GM-CSF, Flat3-ligand), microspheres, dimethyl dioctadecyl ammoniumbromide (DDA) based adjuvants, AS-1 (Smith Kline Beecham), AS-2 (Smith Kline Beecham), Ribi Adjuvant system based adjuvants, QS21 (Aquila), saponin based adjuvants (crude saponin, the saponin Quil A), Syntex adjuvant in its microfluidized form (SAF-m), MV, ddMV (Genesis), immune stimulating complex (iscom) based adjuvants and inactivated toxins.

The compositions and vaccines described herein may be administered as part of a sustained release formulation (*i.e.*, a formulation such as a capsule or sponge that effects a slow release of compound following administration). Such formulations may generally be prepared using well known technology and administered by, for example, oral, rectal or subcutaneous implantation, or by implantation at the desired target site. Sustained-release formulations may contain a polypeptide, polynucleotide, antibody or cell dispersed in a carrier matrix and/or contained within a reservoir surrounded by a rate controlling membrane. Carriers for use within such formulations are biocompatible, and may also be biodegradable; preferably the formulation provides a relatively constant level of active component release. The amount of active compound contained within a sustained release formulation depends upon the site of implantation, the rate and expected duration of release and the nature of the condition to be treated or prevented.

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THERAPY OF MALIGNANT DISEASES

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In further aspects of the present invention, the compositions and vaccines described herein may be used to inhibit the development of malignant diseases (e.g., progressive or metastatic diseases or diseases characterized by small tumor burden such as minimal residual disease). In general, such methods may be used to prevent, delay or treat a disease associated with WT1 expression. In other words, therapeutic methods provided herein may be used to treat an existing WT1-associated disease, or may be used to prevent or delay the onset of such a disease in a patient who is free of disease or who is afflicted with a disease that is not yet associated with WT1 expression.

As used herein, a disease is "associated with WT1 expression" if diseased cells (e.g., tumor cells) at some time during the course of the disease generate detectably higher levels of a WT1 polypeptide than normal cells of the same tissue. Association of WT1 expression with a malignant disease does not require that WT1 be present on a tumor. For example, overexpression of WT1 may be involved with initiation of a tumor, but the protein expression may subsequently be lost. Alternatively, a malignant disease that is not characterized by an increase in WT1 expression may, at a later time, progress to a disease that is characterized by increased WT1 expression. Accordingly, any malignant disease in which diseased cells formerly expressed, currently express or are expected to subsequently express increased levels of WT1 is considered to be "associated with WT1 expression."

Immunotherapy may be performed using any of a variety of techniques, in which compounds or cells provided herein function to remove WT1-expressing cells from a patient. Such removal may take place as a result of enhancing or inducing an immune response in a patient specific for WT1 or a cell expressing WT1. Alternatively, WT1-expressing cells may be removed *ex vivo* (*e.g.*, by treatment of autologous bone marrow, peripheral blood or a fraction of bone marrow or peripheral blood. Fractions of bone marrow or peripheral blood may be obtained using any standard technique in the art.

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Within such methods, pharmaceutical compositions and vaccines may be administered to a patient. As used herein, a "patient" refers to any warm-blooded animal, preferably a human. A patient may or may not be afflicted with a malignant disease. Accordingly, the above pharmaceutical compositions and vaccines may be used to prevent the onset of a disease (i.e., prophylactically) or to treat a patient afflicted with a disease (e.g., to prevent or delay progression and/or metastasis of an existing disease). A patient afflicted with a disease may have a minimal residual disease (e.g., a low tumor burden in a leukemia patient in complete or partial remission or a cancer patient following reduction of the tumor burden after surgery radiotherapy and/or chemotherapy). Such a patient may be immunized to inhibit a relapse (i.e., prevent or delay the relapse, or decrease the severity of a relapse). Within certain preferred embodiments, the patient is afflicted with a leukemia (e.g., AML, CML, ALL or childhood ALL), a myelodysplastic syndrome (MDS) or a cancer (e.g., gastrointestinal, lung, thyroid or breast cancer or a melanoma), where the cancer or leukemia is WT1 positive (i.e., reacts detectably with an anti-WT1 antibody, as provided herein or expresses WT1 mRNA at a level detectable by RT-PCR, as described herein) or suffers from an autoimmune disease directed against WT1-expressing cells.

The compositions provided herein may be used alone or in combination with conventional therapeutic regimens such as surgery, irradiation, chemotherapy and/or bone marrow transplantation (autologous, syngeneic, allogeneic or unrelated). As discussed in greater detail below, binding agents and T cells as provided herein may be used for purging of autologous stem cells. Such purging may be beneficial prior to, for example, bone marrow transplantation or transfusion of blood or components thereof. Binding agents, T cells, antigen presenting cells (APC) and compositions provided herein may further be used for expanding and stimulating (or priming) autologous, allogeneic, syngeneic or unrelated WT1-specific T-cells *in vitro* and/or *in vivo*. Such WT1-specific T cells may be used, for example, within donor lymphocyte infusions.

Routes and frequency of administration, as well as dosage, will vary from individual to individual, and may be readily established using standard techniques. In

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general, the pharmaceutical compositions and vaccines may be administered by injection (e.g., intracutaneous, intramuscular, intravenous or subcutaneous), intranasally (e.g., by aspiration) or orally. In some tumors, pharmaceutical compositions or vaccines may be administered locally (by, for example, rectocoloscopy, gastroscopy, videoendoscopy, angiography or other methods known in the art). Preferably, between 1 and 10 doses may be administered over a 52 week period. Preferably, 6 doses are administered, at intervals of 1 month, and booster vaccinations may be given periodically thereafter. protocols may be appropriate for individual patients. A suitable dose is an amount of a compound that, when administered as described above, is capable of promoting an antitumor immune response that is at least 10-50% above the basal (i.e., untreated) level. Such response can be monitored by measuring the anti-tumor antibodies in a patient or by vaccine-dependent generation of cytolytic effector cells capable of killing the patient's tumor cells in vitro. Such vaccines should also be capable of causing an immune response that leads to an improved clinical outcome (e.g., more frequent complete or partial remissions, or longer disease-free and/or overall survival) in vaccinated patients as compared to non-vaccinated patients. In general, for pharmaceutical compositions and vaccines comprising one or more polypeptides, the amount of each polypeptide present in a dose ranges from about 100 µg to 5 mg. Suitable dose sizes will vary with the size of the patient, but will typically range from about 0.1 mL to about 5 mL.

In general, an appropriate dosage and treatment regimen provides the active compound(s) in an amount sufficient to provide therapeutic and/or prophylactic benefit. Such a response can be monitored by establishing an improved clinical outcome (e.g., more frequent complete or partial remissions, or longer disease-free and/or overall survival) in treated patients as compared to non-treated patients. Increases in preexisting immune responses to WT1 generally correlate with an improved clinical outcome. Such immune responses may generally be evaluated using standard proliferation, cytotoxicity or cytokine assays, which may be performed using samples obtained from a patient before and after treatment.

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Within further aspects, methods for inhibiting the development of a malignant disease associated with WT1 expression involve the administration of autologous T cells that have been activated in response to a WT1 polypeptide or WT1-expressing APC, as described above. Such T cells may be CD4⁺ and/or CD8⁺, and may be proliferated as described above. The T cells may be administered to the individual in an amount effective to inhibit the development of a malignant disease. Typically, about 1×10^9 to 1×10^{11} T cells/M² are administered intravenously, intracavitary or in the bed of a resected tumor. It will be evident to those skilled in the art that the number of cells and the frequency of administration will be dependent upon the response of the patient.

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Within certain embodiments, T cells may be stimulated prior to an autologous bone marrow transplantation. Such stimulation may take place *in vivo* or *in vitro*. For *in vitro* stimulation, bone marrow and/or peripheral blood (or a fraction of bone marrow or peripheral blood) obtained from a patient may be contacted with a WT1 polypeptide, a polynucleotide encoding a WT1 polypeptide and/or an APC that expresses a WT1 polypeptide under conditions and for a time sufficient to permit the stimulation of T cells as described above. Bone marrow, peripheral blood stem cells and/or WT1-specific T cells may then be administered to a patient using standard techniques.

Within related embodiments, T cells of a related or unrelated donor may be stimulated prior to a syngeneic or allogeneic (related or unrelated) bone marrow transplantation. Such stimulation may take place *in vivo* or *in vitro*. For *in vitro* stimulation, bone marrow and/or peripheral blood (or a fraction of bone marrow or peripheral blood) obtained from a related or unrelated donor may be contacted with a WT1 polypeptide, WT1 polynucleotide and/or APC that expresses a WT1 polypeptide under conditions and for a time sufficient to permit the stimulation of T cells as described above. Bone marrow, peripheral blood stem cells and/or WT1-specific T cells may then be administered to a patient using standard techniques.

Within other embodiments, WT1-specific T cells as described herein may be used to remove cells expressing WT1 from autologous bone marrow, peripheral blood or a

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fraction of bone marrow or peripheral blood (e.g., CD34⁺ enriched peripheral blood (PB) prior to administration to a patient). Such methods may be performed by contacting bone marrow or PB with such T cells under conditions and for a time sufficient to permit the reduction of WT1 expressing cells to less than 10%, preferably less than 5% and more preferably less than 1%, of the total number of myeloid or lymphatic cells in the bone marrow or peripheral blood. The extent to which such cells have been removed may be readily determined by standard methods such as, for example, qualitative and quantitative PCR analysis, morphology, immunohistochemistry and FACS analysis. Bone marrow or PB (or a fraction thereof) may then be administered to a patient using standard techniques.

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DIAGNOSTIC METHODS

The present invention further provides methods for detecting a malignant disease associated with WT1 expression, and for monitoring the effectiveness of an immunization or therapy for such a disease. Such methods are based on the discovery, within the present invention, that an immune response specific for WT1 protein can be detected in patients afflicted with such diseases, and that methods which enhance such immune responses may provide a preventive or therapeutic benefit.

To determine the presence or absence of a malignant disease associated with WT1 expression, a patient may be tested for the level of T cells specific for WT1. Within certain methods, a biological sample comprising CD4⁺ and/or CD8⁺ T cells isolated from a patient is incubated with a WT1 polypeptide, a polynucleotide encoding a WT1 polypeptide and/or an APC that expresses a WT1 polypeptide, and the presence or absence of specific activation of the T cells is detected, as described herein. Suitable biological samples include, but are not limited to, isolated T cells. For example, T cells may be isolated from a patient by routine techniques (such as by Ficoll/Hypaque density gradient centrifugation of peripheral blood lymphocytes). T cells may be incubated *in vitro* for 2-9 days (typically 4 days) at 37°C with WT1 polypeptide (e.g., 5 - 25 μg/ml). It may be desirable to incubate another aliquot of a T cell sample in the absence of WT1 polypeptide

to serve as a control. For CD4⁺ T cells, activation is preferably detected by evaluating proliferation of the T cells. For CD8⁺ T cells, activation is preferably detected by evaluating cytolytic activity. A level of proliferation that is at least two fold greater and/or a level of cytolytic activity that is at least 20% greater than in disease-free patients indicates the presence of a malignant disease associated with WT1 expression. Further correlation may be made, using methods well known in the art, between the level of proliferation and/or cytolytic activity and the predicted response to therapy. In particular, patients that display a higher antibody, proliferative and/or lytic response may be expected to show a greater response to therapy.

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Within other methods, a biological sample obtained from a patient is tested for the level of antibody specific for WT1. The biological sample is incubated with a WT1 polypeptide, a polynucleotide encoding a WT1 polypeptide and/or an APC that expresses a WT1 polypeptide under conditions and for a time sufficient to allow immunocomplexes to form. Immunocomplexes formed between the WT1 polypeptide and antibodies in the biological sample that specifically bind to the WT1 polypeptide are then detected. A biological sample for use within such methods may be any sample obtained from a patient that would be expected to contain antibodies. Suitable biological samples include blood, sera, ascites, bone marrow, pleural effusion, and cerebrospinal fluid.

The biological sample is incubated with the WT1 polypeptide in a reaction mixture under conditions and for a time sufficient to permit immunocomplexes to form between the polypeptide and antibodies specific for WT1. For example, a biological sample and WT1 polypeptide may be incubated at 4°C for 24-48 hours.

Following the incubation, the reaction mixture is tested for the presence of immunocomplexes. Detection of immunocomplexes formed between the WT1 polypeptide and antibodies present in the biological sample may be accomplished by a variety of known techniques, such as radioimmunoassays (RIA) and enzyme linked immunosorbent assays (ELISA). Suitable assays are well known in the art and are amply described in the scientific and patent literature (e.g., Harlow and Lane, Antibodies: A Laboratory Manual,

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Cold Spring Harbor Laboratory, 1988). Assays that may be used include, but are not limited to, the double monoclonal antibody sandwich immunoassay technique of David et al. (U.S. Patent 4,376,110); monoclonal-polyclonal antibody sandwich assays (Wide et al., in Kirkham and Hunter, eds., *Radioimmunoassay Methods*, E. and S. Livingstone, Edinburgh, 1970); the "western blot" method of Gordon et al. (U.S. Patent 4,452,901); immunoprecipitation of labeled ligand (Brown et al., *J. Biol. Chem. 255*:4980-4983, 1980); enzyme-linked immunosorbent assays as described by, for example, Raines and Ross (*J. Biol. Chem. 257*:5154-5160, 1982); immunocytochemical techniques, including the use of fluorochromes (Brooks et al., *Clin. Exp. Immunol. 39*: 477, 1980); and neutralization of activity (Bowen-Pope et al., *Proc. Natl. Acad. Sci. USA 81*:2396-2400, 1984). Other immunoassays include, but are not limited to, those described in U.S. Patent Nos.: 3,817,827; 3,850,752; 3,901,654; 3,935,074; 3,984,533; 3,996,345; 4,034,074; and 4,098,876.

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For detection purposes, WT1 polypeptide may either be labeled or unlabeled. Unlabeled WT1 polypeptide may be used in agglutination assays or in combination with labeled detection reagents that bind to the immunocomplexes (e.g., anti-immunoglobulin, protein G, protein A or a lectin and secondary antibodies, or antigen-binding fragments thereof, capable of binding to the antibodies that specifically bind to the WT1 polypeptide). If the WT1 polypeptide is labeled, the reporter group may be any suitable reporter group known in the art, including radioisotopes, fluorescent groups, luminescent groups, enzymes, biotin and dye particles.

Within certain assays, unlabeled WT1 polypeptide is immobilized on a solid support. The solid support may be any material known to those of ordinary skill in the art to which the polypeptide may be attached. For example, the solid support may be a test well in a microtiter plate or a nitrocellulose or other suitable membrane. Alternatively, the support may be a bead or disc, such as glass, fiberglass, latex or a plastic material such as polystyrene or polyvinylchloride. The support may also be a magnetic particle or a fiber optic sensor, such as those disclosed, for example, in U.S. Patent No. 5,359,681. The

polypeptide may be immobilized on the solid support using a variety of techniques known to those of skill in the art, which are amply described in the patent and scientific literature. In the context of the present invention, the term "immobilization" refers to both noncovalent association, such as adsorption, and covalent attachment (which may be a direct linkage between the antigen and functional groups on the support or may be a linkage by way of a cross-linking agent). Immobilization by adsorption to a well in a microtiter plate or to a membrane is preferred. In such cases, adsorption may be achieved by contacting the WT1 polypeptide, in a suitable buffer, with the solid support for a suitable amount of time. The contact time varies with temperature, but is typically between about 1 hour and about 1 day. In general, contacting a well of a plastic microtiter plate (such as polystyrene or polyvinylchloride) with an amount of polypeptide ranging from about 10 ng to about 10 µg, and preferably about 100 ng to about 1 µg, is sufficient to immobilize an adequate amount of polypeptide.

Following immobilization, the remaining protein binding sites on the support are typically blocked. Any suitable blocking agent known to those of ordinary skill in the art, such as bovine serum albumin, Tween 20™ (Sigma Chemical Co., St. Louis, MO), heat-inactivated normal goat serum (NGS), or BLOTTO (buffered solution of nonfat dry milk which also contains a preservative, salts, and an antifoaming agent). The support is then incubated with a biological sample suspected of containing specific antibody. The sample can be applied neat, or, more often, it can be diluted, usually in a buffered solution which contains a small amount (0.1%-5.0% by weight) of protein, such as BSA, NGS, or BLOTTO. In general, an appropriate contact time (*i.e.*, incubation time) is a period of time that is sufficient to detect the presence of antibody that specifically binds WT1 within a sample containing such an antibody. Preferably, the contact time is sufficient to achieve a level of binding that is at least about 95% of that achieved at equilibrium between bound and unbound antibody. Those of ordinary skill in the art will recognize that the time necessary to achieve equilibrium may be readily determined by assaying the level of

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binding that occurs over a period of time. At room temperature, an incubation time of about 30 minutes is generally sufficient.

Unbound sample may then be removed by washing the solid support with an appropriate buffer, such as PBS containing 0.1% Tween 20TM. A detection reagent that binds to the immunocomplexes and that comprises a reporter group may then be added. The detection reagent is incubated with the immunocomplex for an amount of time sufficient to detect the bound antibody. An appropriate amount of time may generally be determined by assaying the level of binding that occurs over a period of time. Unbound detection reagent is then removed and bound detection reagent is detected using the reporter group. The method employed for detecting the reporter group depends upon the nature of the reporter group. For radioactive groups, scintillation counting or autoradiographic methods are generally appropriate. Spectroscopic methods may be used to detect dyes, luminescent groups and fluorescent groups. Biotin may be detected using avidin, coupled to a different reporter group (commonly a radioactive or fluorescent group or an enzyme). Enzyme reporter groups (e.g., horseradish peroxidase, beta-galactosidase, alkaline phosphatase and glucose oxidase) may generally be detected by the addition of substrate (generally for a specific period of time), followed by spectroscopic or other analysis of the reaction products. Regardless of the specific method employed, a level of bound detection reagent that is at least two fold greater than background (i.e., the level observed for a biological sample obtained from a disease-free individual) indicates the presence of a malignant disease associated with WT1 expression.

In general, methods for monitoring the effectiveness of an immunization or therapy involve monitoring changes in the level of antibodies or T cells specific for WT1 in the patient. Methods in which antibody levels are monitored may comprise the steps of:

(a) incubating a first biological sample, obtained from a patient prior to a therapy or immunization, with a WT1 polypeptide, wherein the incubation is performed under conditions and for a time sufficient to allow immunocomplexes to form; (b) detecting immunocomplexes formed between the WT1 polypeptide and antibodies in the biological

sample that specifically bind to the WT1 polypeptide; (c) repeating steps (a) and (b) using a second biological sample taken from the patient following therapy or immunization; and (d) comparing the number of immunocomplexes detected in the first and second biological samples. Alternatively, a polynucleotide encoding a WT1 polypeptide, or an APC expressing a WT1 polypeptide may be employed in place of the WT1 polypeptide. Within such methods, immunocomplexes between the WT1 polypeptide encoded by the polynucleotide, or expressed by the APC, and antibodies in the biological sample are detected.

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Methods in which T cell activation and/or the number of WT1 specific precursors are monitored may comprise the steps of: (a) incubating a first biological sample comprising CD4+ and/or CD8+ cells (e.g., bone marrow, peripheral blood or a fraction thereof), obtained from a patient prior to a therapy or immunization, with a WT1 polypeptide, wherein the incubation is performed under conditions and for a time sufficient to allow specific activation, proliferation and/or lysis of T cells; (b) detecting an amount of activation, proliferation and/or lysis of the T cells; (c) repeating steps (a) and (b) using a second biological sample comprising CD4+ and/or CD8+ T cells, and taken from the same patient following therapy or immunization; and (d) comparing the amount of activation, proliferation and/or lysis of T cells in the first and second biological samples. Alternatively, a polynucleotide encoding a WT1 polypeptide, or an APC expressing a WT1 polypeptide may be employed in place of the WT1 polypeptide.

A biological sample for use within such methods may be any sample obtained from a patient that would be expected to contain antibodies, CD4+ T cells and/or CD8+ T cells. Suitable biological samples include blood, sera, ascites, bone marrow, pleural effusion and cerebrospinal fluid. A first biological sample may be obtained prior to initiation of therapy or immunization or part way through a therapy or vaccination regime. The second biological sample should be obtained in a similar manner, but at a time following additional therapy or immunization. The second biological sample may be obtained at the completion of, or part way through, therapy or immunization, provided that

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at least a portion of therapy or immunization takes place between the isolation of the first and second biological samples.

Incubation and detection steps for both samples may generally be performed as described above. A statistically significant increase in the number of immunocomplexes in the second sample relative to the first sample reflects successful therapy or immunization.

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The following Examples are offered by way of illustration and not by way of limitation.

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EXAMPLES

Example 1

Identification of an Immune Response to WT1

in Patients with Hematological Malignancies

This Example illustrates the identification of an existent immune response in patients with a hematological malignancy.

To evaluate the presence of preexisting WT1 specific antibody responses in patients, sera of patients with AML, ALL, CML and severe aplastic anemia were analyzed using Western blot analysis. Sera were tested for the ability to immunoprecipitate WT1 from the human leukemic cell line K562 (American Type Culture Collection, Manassas, VA). In each case, immunoprecipitates were separated by gel electrophoresis, transferred to membrane and probed with the anti WT-1 antibody WT180 (Santa Cruz Biotechnology, Inc., Santa Cruz, CA). This Western blot analysis identified potential WT1 specific antibodies in patients with hematological malignancy. A representative Western blot showing the results for a patient with AML is shown in Figure 2. A 52 kD protein in the immunoprecipitate generated using the patient sera was recognized by the WT1 specific antibody. The 52 kD protein migrated at the same size as the positive control.

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Example 2

Induction of Antibodies to WT1 in Mice Immunized with Cell Lines Expressing WT1

This Example illustrates the use of cells expressing WT1 to induce a WT1 specific antibody response in vivo.

Detection of existent antibodies to WT1 in patients with leukemia strongly implied that it is possible to immunize to WT1 protein to elicit immunity to WT1. To test whether immunity to WT1 can be generated by vaccination, mice were injected with TRAMP-C, a WT1 positive tumor cell line of B6 origin. Briefly, male B6 mice were

immunized with 5 x 10^6 TRAMP-C cells subcutaneously and boosted twice with 5 x 10^6 cells at three week intervals. Three weeks after the final immunization, sera were obtained and single cell suspensions of spleens were prepared in RPMI 1640 medium (GIBCO) with 25μ M β -2-mercaptoethanol, 200 units of penicillin per ml, 10mM L-glutamine, and 10% fetal bovine serum.

Following immunization to TRAMP-C, a WT1 specific antibody response in the immunized animals was detectable. A representative Western blot is shown in Figure 3. These results show that immunization to WT1 protein can elicit an immune response to WT1 protein.

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Example 3 Induction of Th and Antibody Responses in Mice Immunized with WT1 Peptides

This Example illustrates the ability of immunization with WT1 peptides to elicit an immune response specific for WT1.

Peptides suitable for eliciting Ab and proliferative T cell responses were identified according to the Tsites program (Rothbard and Taylor, *EMBO J. 7*:93-100, 1988; Deavin et al., *Mol. Immunol. 33*:145-155, 1996), which searches for peptide motifs that have the potential to elicit Th responses. Peptides shown in Table I were synthesized and sequenced.

Table I WT1 Peptides

Peptide	Sequence	Comments
Mouse: p6-22	RDLNALLPAVSSLGGGG (SEQ ID NO:13)	1 mismatch relative to human WT1 sequence
Human: p6-22	RDLNALLPAVPSLGGGG (SEQ ID NO:1)	
Human/mouse: p117-139	PSQASSGQARMFPNAPYLPSCLE (SEQ ID NOs: 2 and 3)	

Mouse: p244-262	GATLKGMAAGSSSSVKWTE	1 mismatch relative to
	(SEQ ID NO:14)	human WT1 sequence
Human: p244-262	GATLKGVAAGSSSSVKWTE	
	(SEQ ID NO:4)	
Human/mouse:	RIHTHGVFRGIQDVR	
p287-301	(SEQ ID NOs: 15 and 16)	,
Mouse: p299-313	VRRVSGVAPTLVRS	1 mismatch relative to
	(SEQ ID NO:17)	human WT1 sequence
Human/mouse:	CQKKFARSDELVRHH	
p421-435	(SEQ ID NOs: 19 and 20)	

For immunization, peptides were grouped as follows:

5	Group A:	p6-22 human: 10.9mg in 1ml (10μ l = 100μ g) p117-139 human/mouse: 7.6mg in 1ml (14μ l = 100μ g) p244-262 human: 4.6.mg in 1ml (22μ l = 100μ g)
10	Group B:	p287-301 human/mouse: 7.2mg in 1ml ($14\mu l = 100\mu g$) mouse p299-313: 6.6.mg in 1ml ($15\mu l = 100\mu g$) p421-435 human/mouse: 3.3mg in 1ml ($30\mu l = 100\mu g$)

Control: (FBL peptide 100μg) + CFA/IFA

Control:

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Group A contained peptides present within the amino terminus portion of WT1 (exon 1) and Group B contained peptides present within the carboxy terminus, which contains a four zinc finger region with sequence homology to other DNA-binding proteins. Within group B, p287-301 and p299-313 were derived from exon 7, zinc finger 1, and p421-435 was derived from exon 10, zinc finger IV.

(CD45 peptide 100µg) + CFA/IFA

B6 mice were immunized with a group of WT1 peptides or with a control peptide. Peptides were dissolved in 1ml sterile water for injection, and B6 mice were immunized 3 times at time intervals of three weeks. Adjuvants used were CFA/IFA, GM-CSF, and Montinide. The presence of antibodies specific for WT1 was then determined as

described in Examples 1 and 2, and proliferative T cell responses were evaluated using a standard thymidine incorporation assay, in which cells were cultured in the presence of antigen and proliferation was evaluated by measuring incorporated radioactivity (Chen et al., Cancer Res. 54:1065-1070, 1994). In particular, lymphocytes were cultured in 96-well plates at 2x10⁵ cells per well with 4x10⁵ irradiated (3000 rads) syngeneic spleen cells and the designated peptide.

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Immunization of mice with the group of peptides designated as Group A elicited an antibody response to WT1 (Figure 4). No antibodies were detected following immunization to Vaccine B, which is consistent with a lack of helper T cell response from immunization with Vaccine B. P117-139 elicited proliferative T cell responses (Figures 5A-5C). The stimulation indices (SI) varied between 8 and 72. Other peptides (P6-22 and P299-313) also were shown to elicit proliferative T cell responses. Immunization with P6-22 resulted in a stimulation index (SI) of 2.3 and immunization with P299-313 resulted in a SI of 3.3. Positive controls included ConA stimulated T cells, as well as T cells stimulated with known antigens, such as CD45 and FBL, and allogeneic T cell lines (DeBruijn et al., Eur. J. Immunol. 21:2963-2970, 1991).

Figures 6A and 6B show the proliferative response observed for each of the three peptides within vaccine A (Figure 6A) and vaccine B (Figure 6B). Vaccine A elicited proliferative T cell responses to the immunizing peptides p6-22 and p117-139, with stimulation indices (SI) varying between 3 and 8 (bulk lines). No proliferative response to p244-262 was detected (Figure 6A).

Subsequent *in vitro* stimulations were carried out as single peptide stimulations using only p6-22 and p117-139. Stimulation of the Vaccine A specific T cell line with p117-139 resulted in proliferation to p117-139 with no response to p6-22 (Figure7A). Clones derived from the line were specific for p117-139 (Figure 7B). By contrast, stimulation of the Vaccine A specific T cell line with p6-22 resulted in proliferation to p6-22 with no response to p117-139 (Figure 7C). Clones derived from the line were specific for p6-22 (Figure 7D).

These results show that vaccination with WT1 peptides can elicit antibody responses to WT1 protein and proliferative T cell responses to the immunizing peptides.

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Example 4

Induction of CTL Responses in Mice Immunized with WT1 Peptides

This Example illustrates the ability of WT1 peptides to elicit CTL immunity.

Peptides (9-mers) with motifs appropriate for binding to class I MHC were identified using a BIMAS HLA peptide binding prediction analysis (Parker et al., *J. Immunol. 152*:163, 1994). Peptides identified within such analyses are shown in Tables II - XLIV. In each of these tables, the score reflects the theoretical binding affinity (half-time of dissociation) of the peptide to the MHC molecule indicated.

Peptides identified using the Tsites program (Rothbard and Taylor, *EMBO J.* 7:93-100, 1988; Deavin et al., *Mol. Immunol.* 33:145-155, 1996), which searches for peptide motifs that have the potential to elicit Th responses are further shown in Figures 8A and 8B, and Table XLV.

Table II

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA A1

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	137	CLESQPAIR (SEQ ID	18.000
		NO:47)	
2	80	GAEPHEEQC (SEQ	9.000
		ID NO:87)	
3	40	FAPPGASAY (SEQ	5.000
		ID NO:74)	
4	354	QCDFKDCER (SEQ	5.000
		ID NO:162)	
5	2	GSDVRDLNA (SEQ	3.750

		ID NO:101)	
6	152	VTFDGTPSY (SEQ ID NO:244)	2.500
7	260	WTEGQSNHS (SEQ ID NO:247)	2.250
8	409	TSEKPFSCR (SEQ ID NO:232)	1.350
9	73	KQEPSWGGA (SEQ ID NO:125)	1.350
10	386	KTCQRKFSR (SEQ ID NO:128)	1.250
11	37	VLDFAPPGA (SEQ ID NO:241)	1.000
12	325	CAYPGCNKR (SEQ ID NO:44)	1.000
13	232	QLECMTWNQ (SEQ ID NO:167)	0.900
14	272	ESDNHTTPI (SEQ ID NO:71)	0.750
15	366	RSDQLKRHQ (SEQ ID NO:193)	0.750
16	222	SSDNLYQMT (SEQ ID NO:217)	0.750
17	427	RSDELVRHH (SEQ ID NO:191)	0.750
18	394	RSDHLKTHT (SEQ ID NO:192)	0.750
19	317	TSEKRPFMC (SEQ ID NO:233)	0.675
20	213	QALLLRTPY (SEQ ID NO:160)	0.500

Table III

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA A 0201

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	126	RMFPNAPYL (SEQ	313.968

[ID NO:185)	
2	187	SLGEQQYSV (SEQ	285.163
		ID NO:214)	203.103
3	10	ALLPAVPSL (SEQ ID	181.794
	10	NO:34)	101.751
4	242	NLGATLKGV (SEQ	159.970
		ID NO:146)	
5	225	NLYQMTSQL (SEQ	68.360
		ID NO:147)	
6	292	GVFRGIQDV (SEQ	51.790
	,	ID NO:103)	31.,,30
7	191	QQYSVPPPV (SEQ	22.566
		ID NO:171)	
8	280	ILCGAQYRI (SEQ ID	17.736
		NO:116)	
9	235	CMTWNQMNL (SEQ	15.428
		ID NO:49)	
10	441	NMTKLQLAL (SEQ	15.428
		ID NO:149)	
11	7	DLNALLPAV (SEQ	11.998
		ID NO:58)	
12	227	YQMTSQLEC (SEQ	8.573
		ID NO:251)	
13	239	NQMNLGATL (SEQ	8.014
		ID NO:151)	
14	309	TLVRSASET (SEQ ID	7.452
		NO:226)	
15	408	KTSEKPFSC (SEQ ID	5.743
		NO:129)	
16	340	LQMHSRKHT (SEQ	4.752
		ID NO:139)	
17	228	QMTSQLECM (SEQ	4.044
	-	ID NO:169)	
18	93	TVHFSGQFT (SEQ ID	3.586
 		NO:235)	
19	37	VLDFAPPGA (SEQ	3.378
		ID NO:241)	2.040
20	86	EQCLSAFTV (SEQ ID	3.068
LL	·	NO:69)	

Table IV

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA A 0205

			Score (Estimate of Half Time of
	!	Subsequence Residue	Disassociation of a Molecule Containing
Rank	Start Position	Listing	This Subsequence)
1	10	ALLPAVPSL (SEQ ID	42.000
		NO:34)	
2	292	GVFRGIQDV (SEQ ID	24.000
		NO:103)	
3	126	RMFPNAPYL (SEQ ID	21.000
		NO:185)	
4	225	NLYQMTSQL (SEQ	21.000
		ID NO:147)	
5	239	NQMNLGATL (SEQ	16.800
	•	ID NO:151)	
6	302	RVPGVAPTL (SEQ ID	14.000
		NO:195)	
7	441	NMTKLQLAL (SEQ	7.000
		ID NO:149)	
8	235	CMTWNQMNL (SEQ	7.000
		ID NO:49)	-
9	187	SLGEQQYSV (SEQ ID	6.000
		NO:214)	
10	191	QQYSVPPPV (SEQ ID	4.800
		NO:171)	
11	340	LQMHSRKHT (SEQ	4.080
		ID NO:139)	
12	242	NLGATLKGV (SEQ	4.000
		ID NO:146)	
13	227	YQMTSQLEC (SEQ ID	3.600
		NO:251)	
14	194	SVPPPVYGC (SEQ ID	2.000
<u> </u>		NO:218).	
15	93	TVHFSGQFT (SEQ ID	2.000
		NO:235)	·
16	280	ILCGAQYRI (SEQ ID	1.700
		NO:116)	
17	98	GQFTGTAGA (SEQ ID	1.200
		NO:99)	

18	309	TLVRSASET (SEQ ID NO:226)	1.000
19	81	AEPHEEQCL (SEQ ID NO:30)	0.980
20	73	KQEPSWGGA (SEQ ID NO:125)	0.960

Table V

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA A24

			2 12 12 12
	•		Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	302	RVPGVAPTL (SEQ	16.800
		ID NO:195)	
2	218	RTPYSSDNL (SEQ ID	12.000
		NO:194)	
3	356	DFKDCERRF (SEQ	12.000
1		ID NO:55)	
4	126	RMFPNAPYL (SEQ	9.600
		ID NO:185)	·
5	326	AYPGCNKRY (SEQ	7.500
		ID NO:42)	
6	270	GYESDNHT (SEQ ID	7.500
		NO:106)T	
7	239.	NQMNLGATL (SEQ	7.200
İ i		ID NO:151)	
8	10	ALLPAVPSL (SEQ ID	7.200
1		NO:34)	
9	130	NAPYLPSCL (SEQ ID	7.200
		NO:144)	
10	329	GCNKRYFKL (SEQ	6.600
		ID NO:90)	
11	417	RWPSCQKKF (SEQ	6.600
		ID NO:196)	
12	47	AYGSLGGPA (SEQ	6.000
]		ID NO:41)	
13	180	DPMGQQGSL (SEQ	6.000
		ID NO:59)	

14	4 .	DVRDLNALL (SEQ ID NO:62)	5.760
15	285	QYRIHTHGV (SEQ ID NO:175)	5.000
16	192	QYSVPPPVY (SEQ ID NO:176)	5.000
17	207	DSCTGSQAL (SEQ ID NO:61)	4.800
18	441	NMTKLQLAL (SEQ ID NO:149)	4.800
19	225	NLYQMTSQL (SEQ ID NO:147)	4.000
20	235	CMTWNQMNL (SEQ ID NO:49)	4.000

Table VI

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA A3

		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	436	NMHQRNMTK (SEQ	40.000
		ID NO:148)	
2	240	QMNLGATLK (SEQ	20.000
		ID NO:168)	
3	88	CLSAFTVHF (SEQ ID	6.000
		NO:48)	<u> </u>
4	126	RMFPNAPYL (SEQ	4.500
1		ID NO:185)	
5	169	AQFPNHSFK (SEQ	4.500
	,	ID NO:36)	•
6	10	ALLPAVPSL (SEQ ID	4.050
		NO:34)	
7	137	CLESQPAIR (SEQ ID	4.000
		NO:47)	
8	225	NLYQMTSQL (SEQ	3.000
		ID NO:147)	
9	32	AQWAPVLDF (SEQ	2.700
	<u> </u>	ID NO:37)	

10	280	ILCGAQYRI (SEQ ID	2.700
1		NO:116)	
11	386	KTCQRKFSR (SEQ	1.800
		ID NO:128)	
12	235	CMTWNQMNL (SEQ	1.200
		ID NO:49)	
13	441	NMTKLQLAL (SEQ	1.200
		ID NO:149)	
14	152	VTFDGTPSY (SEQ ID	1.000
	·	NO:244)	
15	187	SLGEQQYSV (SEQ	0.900
		ID NO:214)	
16	383	FQCKTCQRK (SEQ.	0.600
		ID NO:80)	
17	292	GVFRGIQDV (SEQ	0.450
		ID NO:103)	
18	194	SVPPPVYGC (SEQ ID	0.405
		NO:218)	
19	287	RIHTHGVFR (SEQ ID	0.400
		NO:182)	
20	263	GQSNHSTGY (SEQ	0.360
		ID NO:100)	

Table VII

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA A68.1

		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	100	FTGTAGACR (SEQ	100.000
		ID NO:84)	
2	386	KTCQRKFSR (SEQ	50.000
		ID NO:128)	
3	368	DQLKRHQRR (SEQ	30.000
		ID NO:60)	
4	312	RSASETSEK (SEQ ID	18.000
		NO:190)	
. 5	337	LSHLQMHSR (SEQ	15.000
		ID NO:141)	

6	364	FSRSDQLKR (SEQ ID	15.000
	50,	NO:83)	15.500
7	409	TSEKPFSCR (SEQ ID	15.000
	.05	NO:232)	10.000
8	299	DVRRVPGVA (SEQ	12.000
		ID NO:63)	
9	4	DVRDLNALL (SEQ	12.000
		ID NO:62)	
10	118	SQASSGQAR (SEQ	10.000
		ID NO:216)	
11	343	HSRKHTGEK (SEQ	9.000
		ID NO:111)	
12	169	AQFPNHSFK (SEQ	9.000
		ID NO:36)	
13	292	GVFRGIQDV (SEQ	8.000
		ID NO:103)	
14	325	CAYPGCNKR (SEQ	7.500
		ID NO:44)	
15	425	FARSDELVR (SEQ	7.500
		ID NO:75)	
16	354	QCDFKDCER (SEQ	7.500
		ID NO:162)	
17	324	MCAYPGCNK (SEQ	6.000
		ID NO:142)	·
18	251	AAGSSSSVK (SEQ	6.000
		ID NO:28)	
19	379	GVKPFQCKT (SEQ	6.000
		ID NO:104)	
20	137	CLESQPAIR (SEQ ID	5.000
		NO:47)	

<u>Table VIII</u> <u>Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA A 1101</u>

Rank	Start Position	Subsequence Residue Listing	Score (Estimate of Half Time of Disassociation of a Molecule Containing This Subsequence)
1	386	KTCQRKFSR (SEQ ID NO:128)	1.800

2	169	AQFPNHSFK (SEQ ID NO:36)	1.200
3	436	NMHQRNMTK (SEQ	0.800
		ID NO:148)	
4	391	KFSRSDHLK (SEQ	0.600
		ID NO:120)	
5	373	HQRRHTGVK (SEQ	0.600
		ID NO:109)	
6	383	FQCKTCQRK (SEQ	0.600
		ID NO:80)	
7	363	RFSRSDQLK (SEQ ID	0.600
		NO:178)	· · · · · · · · · · · · · · · · · · ·
8	240	QMNLGATLK (SEQ	0.400
		ID NO:168)	
9	287	RIHTHGVFR (SEQ ID	0.240
		NO:182)	
10	100	FTGTAGACR (SEQ	0.200
		ID NO:84)	
11	324	MCAYPGCNK (SEQ	0.200
10		ID NO:142)	
12	251	AAGSSSSVK (SEQ	0.200
		ID NO:28)	
13	415	SCRWPSCQK (SEQ	0.200
		ID NO:201)	
14	118	SQASSGQAR (SEQ	0.120
		ID NO:216)	0.100
15	292	GVFRGIQDV (SEQ	0.120
16	137	ID NO:103)	0.000
16	137	CLESQPAIR (SEQ ID	0.080
17	425	NO:47)	0.000
1/	423	FARSDELVR (SEQ	0.080
18	325	ID NO:75) CAYPGCNKR (SEQ	0.080
10	323	ID NO:44)	U.U6U
19	312	RSASETSEK (SEQ ID	0.060
17	312	NO:190)	0.000
20	65	PPPPHSFI (SEQ ID	0.060
20	05	NO:156)K	0.000
	-1	110.130/1	

Table IX

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA A 3101

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	386	KTCQRKFSR (SEQ	9.000
		ID NO:128)	
2	287	RIHTHGVFR (SEQ ID	6.000
		NO:182)	
3	137	CLESQPAIR (SEQ ID	2.000
		NO:47)	
4	118	SQASSGQAR (SEQ	2.000
	·	ID [,] NO:216)	
5	368	DQLKRHQRR (SEQ	1.200
		ID NO:60)	•
6	100	FTGTAGACR (SEQ	1.000
		ID NO:84)	
7	293	VFRGIQDVR (SEQ ID	0.600
		NO:238)	
8	325	CAYPGCNKR (SEQ	0.600
		ID NO:44)	
9	169	AQFPNHSFK (SEQ	0.600
		ID NO:36)	
10	279	PILCGAQYR (SEQ ID	0.400
		NO:155)	
11	436	NMHQRNMTK (SEQ	0.400
		ID NO:148)	
12	425	FARSDELVR (SEQ	0.400
		ID NO:75)	
13	32	AQWAPVLDF (SEQ	0.240
		ID NO:37)	
14	240	QMNLGATLK (SEQ	0.200
		ID NO:168)	
15	354	QCDFKDCER (SEQ	0.200
		ID NO:162)	
16	373	HQRRHTGVK (SEQ	0.200
		ID NO:109)	
17	383	FQCKTCQRK (SEQ	0.200
	<u> </u>	ID NO:80)	

18	313	SASETSEKR (SEQ ID NO:197)	0.200
19	358	KDCERRFSR (SEQ ID NO:118)	0.180
20	391	KFSRSDHLK (SEQ ID NO:120)	0.180

Table X

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA A 3302

		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	337	LSHLQMHSR (SEQ	15.000
1	337	ID NO:141)	15.000
2	409	TSEKPFSCR (SEQ ID NO:232)	15.000
3	364	FSRSDQLKR (SEQ ID NO:83)	15.000
4	137	CLESQPAIR (SEQ ID NO:47)	9.000
5	368	DQLKRHQRR (SEQ ID NO:60)	9.000
6	287	RIHTHGVFR (SEQ ID NO:182)	4.500
7	210	TGSQALLLR (SEQ ID NO:223)	3.000
8	425	FARSDELVR (SEQ ID NO:75)	3.000
9	313	SASETSEKR (SEQ ID NO:197)	3.000
10	293	VFRGIQDVR (SEQ ID NO:238)	3.000
11	354	QCDFKDCER (SEQ ID NO:162)	3.000
12	100	FTGTAGACR (SEQ ID NO:84)	3.000
13	118	SQASSGQAR (SEQ ID NO:216)	3.000

14	325	CAYPGCNKR (SEQ ID NO:44)	3.000
15	207	DSCTGSQAL (SEQ ID NO:61)	1.500
16	139	ESQPAIRNQ (SEQ ID NO:72)	1.500
17	299	DVRRVPGVA (SEQ ID NO:63)	1.500
18	419	PSCQKKFAR (SEQ ID NO:159)	1.500
19	272	ESDNHTTPI (SEQ ID NO:71)	1.500
20	4	DVRDLNALL (SEQ ID NO:62)	1.500

Table XI

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B14

		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	362	RRFSRSDQL (SEQ ID NO:187)	1000.000
2	332	KRYFKLSHL (SEQ ID NO:127)	300.000
3	423	KKFARSDEL (SEQ ID NO:122)	150.000
4	390	RKFSRSDHL (SEQ ID NO:183)	150.000
5	439	QRNMTKLQL (SEQ ID NO:173)	20.000
6	329	GCNKRYFKL (SEQ ID NO:90)	10.000
7	10	ALLPAVPSL (SEQ ID NO:34)	10.000
8	180	DPMGQQGSL (SEQ ID NO:59)	9.000
9	301	RRVPGVAPT (SEQ ID NO:189)	6.000

10	126	DATEDNIA DVI (CEO	5 000
10	126	RMFPNAPYL (SEQ	5.000
		ID NO:185)	
11	371	KRHQRRHTG (SEQ	5.000
		ID NO:126)	
12	225	NLYQMTSQL (SEQ	5.000
		ID NO:147)	
13	144	IRNQGYSTV (SEQ ID	4.000
1		NO:117)	
14	429	DELVRHHNM (SEQ	3.000
		ID NO:53)	
15	437	MHQRNMTKL (SEQ	3.000
		ID NO:143)	
16	125	ARMFPNAPY (SEQ	3.000
		ID NO:38)	•
17	239	NQMNLGATL (SEQ	3.000
		ID NO:151)	,
18	286	YRIHTHGVF (SEQ ID	3.000
		NO:252)	
19	174	HSFKHEDPM (SEQ	3.000
]		ID NO:110)	
20	372	RHQRRHTGV (SEQ	3.000
1		ID NO:181)	

Table XII

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B40

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	81	AEPHEEQCL (SEQ ID	40.000
		NO:30)	
2	429	DELVRHHNM (SEQ	24.000
		ID NO:53)	
3	410	SEKPFSCRW (SEQ	20.000
		ID NO:207)	
4	318	SEKRPFMCA (SEQ	15.000
		ID NO:208)	
5	233	LECMTWNQM (SEQ	12.000
1		ID NO:131)	

6	3	SDVRDLNAL (SEQ	10.000
	,	ID NO:206)	10.000
7	349	GEKPYQCDF (SEQ	8.000
'	3.7	ID NO:91)	0.000
8	6	RDLNALLPA (SEQ	5.000
	-	ID NO:177)	2.000
9	85	EEQCLSAFT (SEQ ID	4.000
		NO:65)	
10	315	SETSEKRPF (SEQ ID	4.000
		NO:209)	
11	261	TEGQSNHST (SEQ ID	4.000
		NO:221)	
12	23	GCALPVSGA (SEQ	3.000
	•	ID NO:89)	
13	38	LDFAPPGAS (SEQ ID	3.000
		NO:130)	
14	273	SDNHTTPIL (SEQ ID	2.500
		NO:204)	
15	206	TDSCTGSQA (SEQ	2.500
		ID NO:220)	
16	24	CALPVSGAA (SEQ	2.000
		ID NO:43)	
17	98	GQFTGTAGA (SEQ	2.000
	······································	ID NO:99)	
18	30	GAAQWAPVL (SEQ	2.000
		ID NO:86)	
19	84	HEEQCLSAF (SEQ ID	2.000
		NO:107)	0.000
20	26	LPVSGAAQW (SEQ	2.000
		ID NO:138)	

Table XIII

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B60

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	81	AEPHEEQCL (SEQ ID	160.000
		NO:30)	

2	3	SDVRDLNAL (SEQ	40.000
		ID NO:206)	
3	429	DELVRHHNM (SEQ	40.000
		ID NO:53)	
4	233	LECMTWNQM (SEQ	22.000
		ID NO:131)	
5	273	SDNHTTPIL (SEQ ID	20.000
	2,5	NO:204)	20.000
6	209	CTGSQALLL (SEQ ID	8.000
		NO:52)	
7	30	GAAQWAPVL (SEQ	8.000
		ID NO:86)	
8	318	SEKRPFMCA (SEQ	8.000
		ID NO:208)	
9	180	DPMGQQGSL (SEQ	8.000
]		ID NO:59)	
10	138	LESQPAIRN (SEQ ID	5.280
		NO:132)	
11	239	NQMNLGATL (SEQ	4.400
		ID NO:151)	·
12	329	GCNKRYFKL (SEQ	4.400
1		ID NO:90)	
13	130	NAPYLPSCL (SEQ ID	4.400
l i		NO:144)	
14	85	EEQCLSAFT (SEQ ID	4.400
		NO:65)	
15	208	SCTGSQALL (SEQ ID	4.000
		NO:202)	
16	207	DSCTGSQAL (SEQ	4.000
		ID NO:61)	
17	218	RTPYSSDNL (SEQ ID	4.000
		NO:194)	
18	261	TEGQSNHST (SEQ ID	4.000
		NO:221)	
19	18	LGGGGGCAL (SEQ	4.000
		ID NO:134)	
20	221	YSSDNLYQM (SEQ	2.200
		ID NO:253)	

Table XIV

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B61

		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
l	318	SEKRPFMCA (SEQ	20.000
] 310	ID NO:208)	20.000
2	429	DELVRHHNM (SEQ	16.000
2	423	ID NO:53)	10.000
3	298	QDVRRVPGV (SEQ	10.000
3	296	ID NO:164)	10.000
4	81	AEPHEEQCL (SEQ ID	8.000
4	01	NO:30)	8.000
5	233	LECMTWNQM (SEQ	8.000
3	233	ID NO:131)	8.000
6	6	RDLNALLPA (SEQ	5.500
O	l °	ID NO:177)	3.300
7	85	EEQCLSAFT (SEQ ID	4.000
,	. 63	NO:65)	4.000
8	261	TEGQSNHST (SEQ ID	4.000
0	201	NO:221)	4.000
9	206	TDSCTGSQA (SEQ	2.500
	200	ID NO:220)	2.500
10	295	RGIQDVRRV (SEQ	2.200
10	2,5	ID NO:179)	2.200
11	3	SDVRDLNAL (SEQ	2.000
	J	ID NO:206)	2.000
12	250	VAAGSSSSV (SEQ	2.000
		ID NO:236)	2.000
13	29	SGAAQWAPV (SEQ	2.000
		ID NO:211)	2.555
14	315	SETSEKRPF (SEQ ID	1.600
- •		NO:209)	
15	138	LESQPAIRN (SEQ ID	1.200
10	-50	NO:132)	
16	244	GATLKGVAA (SEQ	1.100
•	1	ID NO:88)	
17	20	GGGCALPV (SEQ	1.100
•		ID NO:92)	

18	440	RNMTKLQLA (SEQ ID NO:186)	1.100
19	23	GCALPVSGA (SEQ ID NO:89)	1.100
20	191	QQYSVPPPV (SEQ ID NO:171)	1.000

Table XV

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA B62

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	146	NQGYSTVTF (SEQ	211.200
		ID NO:150)	
2	32	AQWAPVLDF (SEQ	96.000
		ID NO:37)	
3	263	GQSNHSTGY (SEQ	96.000
		ID NO:100)	
4	88	CLSAFTVHF (SEQ ID	96.000
		NO:48)	
5	17	SLGGGGGCA (SEQ	9.600
	· · · · · · · · · · · · · · · · · · ·	ID NO:215)	·
6	239	NQMNLGATL (SEQ	8.800
		ID NO:151)	
7	191	QQYSVPPPV (SEQ	8.000
		ID NO:171)	
8	98	GQFTGTAGA (SEQ	8.000
		ID NO:99)	
9	384	QCKTCQRKF (SEQ	6.000
		ID NO:163)	
10	40	FAPPGASAY (SEQ	4.800
		ID NO:74)	
11	227	YQMTSQLEC (SEQ	4.800
		ID NO:251)	
12	187	SLGEQQYSV (SEQ	4.400
		ID NO:214)	
13	86	EQCLSAFTV (SEQ ID	4.400
		NO:69)	

14	152	VTFDGTPSY (SEQ ID NO:244)	4.400
15	101	TGTAGACRY (SEQ ID NO:224)	4.000
16	242	NLGATLKGV (SEQ ID NO:146)	4.000
17	92	FTVHFSGQF (SEQ ID NO:85)	4.000
18	7	DLNALLPAV (SEQ ID NO:58)	4.000
19	123	GQARMFPNA (SEQ ID NO:98)	4.000
20	280	ILCGAQYRI (SEQ ID NO:116)	3.120

Table XVI
Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B7

	1		Consul (Estimate of Half Time of
			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	180	DPMGQQGSL (SEQ	240.000
		ID NO:59)	
2	4	DVRDLNALL (SEQ	200.000
		ID NO:62)	
3	302	RVPGVAPTL (SEQ	20.000
i	ı	ID NO:195)	•
4	30	GAAQWAPVL (SEQ	12.000
		ID NO:86)	
5	239	NQMNLGATL (SEQ	12.000
		ID NO:151)	
6	130	NAPYLPSCL (SEQ ID	12.000
		NO:144)	
7	10	ALLPAVPSL (SEQ ID	12.000
		NO:34)	
. 8	299	DVRRVPGVA (SEQ	5.000
		ID NO:63)	
9	208	SCTGSQALL (SEQ ID	4.000
		NO:202)	

10 1	202	TANCAL DOWN ALL CORC	4.000
10	303	VPGVAPTLV (SEQ	4.000
		ID NO:242)	
11	18	LGGGGGCAL (SEQ	4.000
		ID NO:134)	•
12	218	RTPYSSDNL (SEQ ID	4.000
		NO:194)	
13	207	DSCTGSQAL (SEQ	4.000
i		ID NO:61)	
14	209	CTGSQALLL (SEQ ID	4.000
		NO:52)	
15	329	GCNKRYFKL (SEQ	4.000
		ID NO:90)	
16	235	CMTWNQMNL (SEQ	4.000
-		ID NO:49)	
17	441	NMTKLQLAL (SEQ	4.000
}		ID NO:149)	
18	126	RMFPNAPYL (SEQ	4.000
· j		ID NO:185)	
19	225	NLYQMTSQL (SEQ	4.000
		ID NO:147)	
20	143	AIRNQGYST (SEQ ID	3.000
		NO:33)	

Table XVII

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B8

			Score (Estimate of Half Time of
1		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	329	GCNKRYFKL (SEQ	16.000
		ID NO:90)	
2	4	DVRDLNALL (SEQ	12.000
		ID NO:62)	
3	316	ETSEKRPFM (SEQ ID	3.000
		NO:73)	
4	180	DPMGQQGSL (SEQ	1.600
		ID NO:59)	
5	208	SCTGSQALL (SEQ ID	0.800
		NO:202)	

6	130	NAPYLPSCL (SEQ ID	0.800
		NO:144)	
7	244	GATLKGVAA (SEQ	0.800
		ID NO:88)	
8	30	GAAQWAPVL (SEQ	0.800
	•	ID NO:86)	
9	299	DVRRVPGVA (SEQ	0.400
		ID NO:63)	
10	420	SCQKKFARS (SEQ	0.400
		ID NO:200)	
11	387	TCQRKFSRS (SEQ ID	0.400
		NO:219)	
12	225	NLYQMTSQL (SEQ	0.400
		ID NO:147)	
13	141	QPAIRNQGY (SEQ	0.400
		ID NO:170)	
14	10	ALLPAVPSL (SEQ ID	0.400
		NO:34)	
15	207	DSCTGSQAL (SEQ	0.400
		ID NO:61)	
16	384	QCKTCQRKF (SEQ	0.400
		ID NO:163)	
17	136	SCLESQPAI (SEQ ID	0.300
		NO:198)	
18	347	HTGEKPYQC (SEQ	0.300
	·	ID NO:112)	
19	401	HTRTHTGKT (SEQ	0.200
		ID NO:114)	
20	332	KRYFKLSHL (SEQ	0.200
		ID NO:127)	

Table XVIII Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA B 2702

Rank	Start Position	Subsequence Residue Listing	Score (Estimate of Half Time of Disassociation of a Molecule Containing This Subsequence)
1	332	KRYFKLSHL (SEQ ID NO:127)	900.000

2	362	RRFSRSDQL (SEQ ID NO:187)	900.000
3	286	YRIHTHGVF (SEQ ID NO:252)	200.000
4	125	ARMFPNAPY (SEQ ID NO:38)	200.000
5	375	RRHTGVKPF (SEQ ID NO:188)	180.000
6	32	AQWAPVLDF (SEQ ID NO:37)	100.000
7	301	RRVPGVAPT (SEQ ID NO:189)	60.000
8	439	QRNMTKLQL (SEQ ID NO:173)	60.000
9	126	RMFPNAPYL (SEQ ID NO:185)	22.500
10	426	ARSDELVRH (SEQ ID NO:39)	20.000
11	146	NQGYSTVTF (SEQ ID NO:150)	20.000
12	144	IRNQGYSTV (SEQ ID NO:117)	20.000
13	389	QRKFSRSDH (SEQ ID NO:172)	20.000
14	263	GQSNHSTGY (SEQ ID NO:100)	20.000
15	416	CRWPSCQKK (SEQ ID NO:50)	20.000
16	191	QQYSVPPPV (SEQ ID NO:171)	10.000
17	217	LRTPYSSDN (SEQ ID NO:140)	10.000
18	107	CRYGPFGPP (SEQ ID NO:51)	10.000
19	98	GQFTGTAGA (SEQ ID NO:99)	10.000
20	239	NQMNLGATL (SEQ ID NO:151)	6.000

Table XIX
Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B 2705

			Score (Estimate of Half Time of
	,	Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	332	KRYFKLSHL (SEQ	30000.000
		ID NO:127)	
2	362	RRFSRSDQL (SEQ ID	30000.000
		NO:187)	•
3	416	CRWPSCQKK (SEQ	10000.000
		ID NO:50)	
4	439	QRNMTKLQL (SEQ	2000.000
		ID NO:173)	į
5	286	YRIHTHGVF (SEQ ID	1000.000
İ		NO:252)	
6	125	ARMFPNAPY (SEQ	1000.000
		ID NO:38)	
7	294	FRGIQDVRR (SEQ ID	1000.000
		NO:81)	
8	432	VRHHNMHQR (SEQ	1000.000
ľ		ID NO:243)	
9	169	AQFPNHSFK (SEQ	1000.000
		ID NO:36)	
10	375	RRHTGVKPF (SEQ	900.000
		ID NO:188)	
11	126	RMFPNAPYL (SEQ	750.000
		ID NO:185)	
12	144	IRNQGYSTV (SEQ ID	600.000
		NO:117)	
13	301	RRVPGVAPT (SEQ	600.000
ļ		¹ ID NO:189)	
14	32	AQWAPVLDF (SEQ	500.000
		ID NO:37)	
15	191	QQYSVPPPV (SEQ	300.000
		ID NO:171)	
16	373	HQRRHTGVK (SEQ	200.000
L		ID NO:109)	
17	426	ARSDELVRH (SEQ	200.000
		ID NO:39)	

18	383	FQCKTCQRK (SEQ ID NO:80)	200.000
19	239	NQMNLGATL (SEQ ID NO:151)	200.000
20	389	QRKFSRSDH (SEQ ID NO:172)	200.000

Table XX

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA B 3501

		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	278	TPILCGAQY (SEQ ID NO:227)	40.000
2	141	QPAIRNQGY (SEQ ID NO:170)	40.000
3	219	TPYSSDNLY (SEQ ID NO:231)	40.000
4	327	YPGCNKRYF (SEQ ID NO:250)	20.000
5	163	TPSHHAAQF (SEQ ID NO:228)	20.000
6	180	DPMGQQGSL (SEQ ID NO:59)	20.000
7	221	YSSDNLYQM (SEQ ID NO:253)	20.000
8	26	LPVSGAAQW (SEQ ID NO:138)	10.000
9	174	HSFKHEDPM (SEQ ID NO:110)	10.000
10	82	EPHEEQCLS (SEQ ID NO:68)	6.000
11	213	QALLLRTPY (SEQ ID NO:160)	6.000
12	119	QASSGQARM (SEQ ID NO:161)	6.000
13	4	DVRDLNALL (SEQ ID NO:62)	6.000

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14	40	FAPPGASAY (SEQ ID NO:74)	6.000
15	120	ASSGQARMF (SEQ ID NO:40)	5.000
16	207	DSCTGSQAL (SEQ ID NO:61)	5.000
17	303	VPGVAPTLV (SEQ ID NO:242)	4.000
18	316	ETSEKRPFM (SEQ ID NO:73)	4.000
19	152	VTFDGTPSY (SEQ ID NO:244)	4.000
20	412	KPFSCRWPS (SEQ ID NO:123)	4.000

Table XXI

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B 3701

		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	3	SDVRDLNAL (SEQ	40.000
	-	ID NO:206)	
2	273	SDNHTTPIL (SEQ ID	40.000
		NO:204)	
3	81	AEPHEEQCL (SEQ ID	10.000
		NO:30)	
4	298	QDVRRVPGV (SEQ	8.000
		ID NO:164)	
5	428	SDELVRHHN (SEQ	6.000
		ID NO:203)	
6	85	EEQCLSAFT (SEQ ID	5.000
		NO:65)	
7	208	SCTGSQALL (SEQ ID	5.000
		NO:202)	
8	4	DVRDLNALL (SEQ	5.000
		ID NO:62)	
9	209	CTGSQALLL (SEQ ID	5.000
		NO:52)	

10	38	LDFAPPGAS (SEQ ID	4.000
10	36	NO:130)	4.000
11	222		4.000
11	223	SDNLYQMTS (SEQ	4.000
		ID NO:205)	
12	179	EDPMGQQGS (SEQ	4.000
		ID NO:64)	
13	206	TDSCTGSQA (SEQ	4.000
		ID NO:220)	
14	6	RDLNALLPA (SEQ	4.000
		ID NO:177)	
15	84	HEEQCLSAF (SEQ ID	2.000
		NO:107)	
16	233	LECMTWNQM (SEQ	2.000
		ID NO:131)	
17	429	DELVRHHNM (SEQ	2.000
		ID NO:53)	
18	315	SETSEKRPF (SEQ ID	2.000
		NO:209)	
19	349	GEKPYQCDF (SEQ	2.000
		ID NO:91)	
20	302	RVPGVAPTL (SEQ	1.500
] ·		ID NO:195)	

Table XXII

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B 3801

Rank	Start Position	Subsequence Residue Listing	Score (Estimate of Half Time of Disassociation of a Molecule Containing This Subsequence)
1	437	MHQRNMTKL (SEQ ID NO:143)	36.000
2	434	HHNMHQRNM (SEQ ID NO:108)	6.000
3	372	RHQRRHTGV (SEQ ID NO:181)	6.000
4	180	DPMGQQGSL (SEQ ID NO:59)	4.000
5	433	RHHNMHQRN (SEQ ID NO:180)	3.900

6	165	SHHAAQFPN (SEQ	3,900
°	103	1	3.900
<u> </u>	202	ID NO:213)	
7	202	CHTPTDSCT (SEQ ID	3.000
		NO:45)	
8	396	DHLKTHTRT (SEQ	3.000
		ID NO:57)	
9	161	GHTPSHHAA (SEQ	3.000
		ID NO:94)	
10	302	RVPGVAPTL (SEQ	2.600
		ID NO:195)	
11	417	RWPSCQKKF (SEQ	2.400
]		ID NO:196)	
12	327	YPGCNKRYF (SEQ	2.400
		ID NO:250)	
13	208	SCTGSQALL (SEQ ID	2.000
		NO:202)	
14	163	TPSHHAAQF (SEQ	2.000
<u> </u>		ID NO:228)	
15	120	ASSGQARMF (SEQ	2.000
		ID NO:40)	
16	18	LGGGGGCAL (SEQ	2.000
		ID NO:134)	·
17	177	KHEDPMGQQ (SEQ	1.800
1		ID NO:121)	
18	83	PHEEQCLSA (SEQ ID	1.800
	•	NO:154)	
19	10	ALLPAVPSL (SEQ ID	1.300
		NO:34)	
20	225	NLYQMTSQL (SEQ	1.300
		ID NO:147)	

Table XXIII Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA B 3901

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	437	MHQRNMTKL (SEQ	135.000
		ID NO:143)	

			· · · · · · · · · · · · · · · · · · ·
2	332	KRYFKLSHL (SEQ ID NO:127)	45.000
3	434	HHNMHQRNM (SEQ ID NO:108)	30.000
4	362	RRFSRSDQL (SEQ ID NO:187)	30.000
5	372	RHQRRHTGV (SEQ ID NO:181)	30.000
6	10	ALLPAVPSL (SEQ ID NO:34)	9.000
7	439	QRNMTKLQL (SEQ ID NO:173)	7.500
8	390	RKFSRSDHL (SEQ ID NO:183)	6.000
9	396	DHLKTHTRT (SEQ ID NO:57)	6.000
10	239	NQMNLGATL (SEQ ID NO:151)	6.000
11	423	KKFARSDEL (SEQ ID NO:122)	6.000
12	126	RMFPNAPYL (SEQ ID NO:185)	6.000
13	225	NLYQMTSQL (SEQ ID NO:147)	6.000
14	180	DPMGQQGSL (SEQ ID NO:59)	6.000
15	144	IRNQGYSTV (SEQ ID NO:117)	5.000
16	136	SCLESQPAI (SEQ ID NO:198)	4.000
17	292	GVFRGIQDV (SEQ ID NO:103)	3.000
18	302	RVPGVAPTL (SEQ ID NO:195)	3.000
19	208	SCTGSQALL (SEQ ID NO:202)	3.000
20	207.	DSCTGSQAL (SEQ ID NO:61)	3.000

Table XXIV

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B 3902

_			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	239	NQMNLGATL (SEQ	24.000
		ID NO:151)	
2	390	RKFSRSDHL (SEQ ID	20.000
		NO:183)	
3	423	KKFARSDEL (SEQ	20.000
		ID NO:122)	
4	32	AQWAPVLDF (SEQ	5.000
		ID NO:37)	
5	146	NQGYSTVTF (SEQ	5.000
		ID NO:150)	
6	130	NAPYLPSCL (SEQ ID	2.400
		NO:144)	
7	225	NLYQMTSQL (SEQ	2.400
		ID NO:147)	
8	30	GAAQWAPVL (SEQ	2.400
		ID NO:86)	
9	441	NMTKLQLAL (SEQ	2.400
		ID NO:149)	
10	302	RVPGVAPTL (SEQ	2.400
		ID NO:195)	
11	126	RMFPNAPYL (SEQ	2.000
<u> </u>		ID NO:185)	
12	218	RTPYSSDNL (SEQ ID	2.000
	200	NO:194)	2.000
13	209	CTGSQALLL (SEQ ID	2.000
1.4	222	NO:52)	2.000
14	332	KRYFKLSHL (SEQ	2.000
1.5	. 100	ID NO:127)	2.000
15	180	DPMGQQGSL (SEQ	2.000
15	427	ID NO:59)	2.000
16	437	MHQRNMTKL (SEQ	2.000
17	207	ID NO:143)	2.000
17	207	DSCTGSQAL (SEQ	2.000
	L	ID NO:61)	

18	208	SCTGSQALL (SEQ ID NO:202)	2.000
19	329	GCNKRYFKL (SEQ ID NO:90)	2.000
20	10	ALLPAVPSL (SEQ ID NO:34)	2.000

Table XXV

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA B 4403

		r	0 (7) 1 (7) (7)
			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	315	SETSEKRPF (SEQ ID	80.000
		NO:209)	
2	349	GEKPYQCDF (SEQ	80.000
		ID NO:91)	
3	84	HEEQCLSAF (SEQ ID	60.000
		NO:107)	
4	410	SEKPFSCRW (SEQ	48.000
	<u> </u>	ID NO:207)	
5	· 429	DELVRHHNM (SEQ	24.000
		ID NO:53)	
6	278	TPILCGAQY (SEQ ID	15.000
		NO:227)	
7	141	QPAIRNQGY (SEQ	9.000
		ID NO:170)	
8	40	FAPPGASAY (SEQ	9.000
		ID NO:74)	
9	213	QALLLRTPY (SEQ ID	9.000
		NO:160)	
10	318	SEKRPFMCA (SEQ	8.000
	ļ	ID NO:208)	
11	81	AEPHEEQCL (SEQ ID	8.000
		NO:30)	
12	152	VTFDGTPSY (SEQ ID	4.500
		NO:244)	
13	101	TGTAGACRY (SEQ	4.500
		ID NO:224)	

14	120	ASSGQARMF (SEQ ID NO:40)	4.500
15	261	TEGQSNHST (SEQ ID NO:221)	4.000
16	85	EEQCLSAFT (SEQ ID NO:65)	4.000
17	233	LECMTWNQM (SEQ ID NO:131)	4.000
18	104	AGACRYGPF (SEQ ID NO:31)	4.000
19	3	SDVRDLNAL (SEQ ID NO:206)	3.000
20	185	QGSLGEQQY (SEQ ID NO:166)	3.000

Table XXVI
Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B 5101

		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
D 1-	Cana Danisian	· •	
Rank	Start Position	Listing	Containing This Subsequence)
1	303	VPGVAPTLV (SEQ	314.600
Į.		ID NO:242)	
2	180	DPMGQQGSL (SEQ	242.000
		ID NO:59)	
. 3	250	VAAGSSSSV (SEQ	157.300
		ID NO:236)	
4	130	NAPYLPSCL (SEQ ID	50.000
	_	NO:144)	
5	30	GAAQWAPVL (SEQ	50.000
		ID NO:86)	
6	20	GGGGCALPV (SEQ	44.000
·		ID NO:92)	
7	64	PPPPPHSFI (SEQ ID	40.000
		NO:157)	
8	29	SGAAQWAPV (SEQ	40.000
<u></u>		ID NO:211)	
9	18	LGGGGGCAL (SEQ	31.460
		ID NO:134)	·

10	295	RGIQDVRRV (SEQ	22.000
		ID NO:179)	22.000
11	119	QASSGQARM (SEQ	18.150
		ID NO:161)	
12	418	WPSCQKKFA (SEQ	12.100
		ID NO:246)	
13	82	EPHEEQCLS (SEQ ID	12.100
		NO:68)	
14	110	GPFGPPPPS (SEQ ID	11.000
		NO:96)	
15	272	ESDNHTTPI (SEQ ID	8.000
		NO:71)	
16	306	VAPTLVRSA (SEQ	7.150
		ID NO:237)	
17	280	ILCGAQYRI (SEQ ID	6.921
		NO:116)	
18	219	TPYSSDNLY (SEQ ID	6.600
		NO:231)	
19	128	FPNAPYLPS (SEQ ID	6.500
		NO:79)	
20	204	TPTDSCTGS (SEQ ID	6.050
		NO:230)	

Table XXVII

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA B 5102

		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	295	RGIQDVRRV (SEQ	290.400
		ID NO:179)	
2	303	VPGVAPTLV (SEQ	200.000
		ID NO:242)	
3	180	DPMGQQGSL (SEQ	133.100
		ID NO:59)	
4	250	VAAGSSSSV (SEQ	110.000
		ID NO:236)	
5	30	GAAQWAPVL (SEQ	55.000
		ID NO:86)	

6	130	NAPYLPSCL (SEQ ID	50.000
"	130	NO:144)	30.000
			44.000
7	20	GGGGCALPV (SEQ	44.000
		ID NO:92)	
8	29	SGAAQWAPV (SEQ	44.000
		ID NO:211)	
9	64	PPPPHSFI (SEQ ID	40.000
		NO:157)	
10	119	QASSGQARM (SEQ	36.300
		ID NO:161)	
11	110	GPFGPPPPS (SEQ ID	27.500
		NO:96)	
12	412	KPFSCRWPS (SEQ ID	25.000
		NO:123)	
13	18	LGGGGCAL (SEQ	24.200
		ID NO:134)	
14	24	CALPVSGAA (SEQ	16.500
		ID NO:43)	
15	219	TPYSSDNLY (SEQ ID	15.000
		NO:231)	
16	292	GVFRGIQDV (SEQ	14.641
		ID NO:103)	
17	136	SCLESQPAI (SEQ ID	14.520
		NO:198)	
18	418	WPSCQKKFA (SEQ	12.100
		ID NO:246)	
19	269	TGYESDNHT (SEQ	11.000
		ID NO:225)	
20	351	KPYQCDFKD (SEQ	11.000
		ID NO:124)	

Table XXVIII Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA B 5201

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		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	191	QQYSVPPPV (SEQ ID NO:171)	100.000

2	32	AQWAPVLDF (SEQ ID NO:37)	30.000
3	243	LGATLKGVA (SEQ ID NO:133)	16.500
4	303	VPGVAPTLV (SEQ ID NO:242)	13.500
5	86	EQCLSAFTV (SEQ ID NO:69)	12.000
6	295	RGIQDVRRV (SEQ ID NO:179)	10.000
7	98	GQFTGTAGA (SEQ ID NO:99)	8.250
8	292	GVFRGIQDV (SEQ ID NO:103)	8.250
9	29	SGAAQWAPV (SEQ ID NO:211)	6.000
10	146	NQGYSTVTF (SEQ ID NO:150)	5.500
11	20	GGGGCALPV (SEQ ID NO:92)	5.000
12	239	NQMNLGATL (SEQ ID NO:151)	4.000
13	64	PPPPPHSFI (SEQ ID NO:157)	3.600
14	273	SDNHTTPIL (SEQ ID NO:204)	3.300
15	286	YRIHTHGVF (SEQ ID NO:252)	3.000
16	269	TGYESDNHT (SEQ ID NO:225)	3.000
17	406	TGKTSEKPF (SEQ ID NO:222)	2.750
18	327	YPGCNKRYF (SEQ ID NO:250)	2.750
19	7	DLNALLPAV (SEQ ID NO:58)	2.640
20	104	AGACRYGPF (SEQ ID NO:31)	2.500

Table XXIX

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA B 5801

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	230	TSQLECMTW (SEQ	96.800
		ID NO:234)	
2	92	FTVHFSGQF (SEQ ID	60.000
		NO:85)	
3	120	ASSGQARMF (SEQ	40.000
		ID NO:40)	
4	168	AAQFPNHSF (SEQ	20.000
		ID NO:29)	
5	408	KTSEKPFSC (SEQ ID	12.000
		NO:129)	
6	394	RSDHLKTHT (SEQ	9.900
		ID NO:192)	
7	276	HTTPILCGA (SEQ ID	7.200
		NO:115)	
8	218	RTPYSSDNL (SEQ ID	6.600
		NO:194)	
9	152	VTFDGTPSY (SEQ ID	6.000
		NO:244)	
10	40	FAPPGASAY (SEQ	6.000
		ID NO:74)	
11	213	QALLLRTPY (SEQ ID	4.500
		NO:160)	
12	347	HTGEKPYQC (SEQ	4.400
		ID NO:112)	
13	252	AGSSSSVKW (SEQ	4.400
1		ID NO:32)	
14	211	GSQALLLRT (SEQ ID	4.356
İ		NO:102)	
15	174	HSFKHEDPM (SEQ	4.000
		ID NO:110)	
16	317	TSEKRPFMC (SEQ	4.000
		ID NO:233)	
17	26	LPVSGAAQW (SEQ	4.000
		ID NO:138)	

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18	289	HTHGVFRGI (SEQ ID NO:113)	3.600
19	222	SSDNLYQMT (SEQ ID NO:217)	3.300
20	96	FSGQFTGTA (SEQ ID NO:82)	3.300

Table XXX

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA CW0301

Score (Estimate of Half Time of Subsequence Residue Disassociation of a Molecule **Start Position** Listing Containing This Subsequence) Rank ALLPAVPSL (SEQ ID 100.000 10 NO:34) KRYFKLSHL (SEQ 2 332 48.000 ID NO:127) RMFPNAPYL (SEQ 3 126 36.000 ID NO:185) 4 3 SDVRDLNAL (SEQ 30.000 ID NO:206) NQMNLGATL (SEQ 5 239 24.000 ID NO:151) NLYQMTSQL (SEQ 6 225 24.000 ID NO:147) 7 180 DPMGQQGSL (SEQ 20.000 ID NO:59) 8 362 RRFSRSDQL (SEQ ID 12.000 NO:187) 9 329 GCNKRYFKL (SEQ 10.000 ID NO:90) YRIHTHGVF (SEQ ID 10 286 10.000 NO:252) 11 301 RRVPGVAPT (SEQ 10.000 ID NO:189) CALPVSGAA (SEQ 12 24 10.000 ID NO:43) SCLESQPAI (SEQ ID 13 136 7.500 NO:198) MHQRNMTKL (SEQ 14 437 7.200

		ID NO:143)	
15	390	RKFSRSDHL (SEQ ID NO:183)	6.000
16	423	KKFARSDEL (SEQ ID NO:122)	6.000
17	92	FTVHFSGQF (SEQ ID NO:85)	5.000
18	429	DELVRHHNM (SEQ ID NO:53)	5.000
19	130	NAPYLPSCL (SEQ ID NO:144)	4.800
20	30	GAAQWAPVL (SEQ ID NO:86)	4.000

Table XXXI
Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Human HLA CW0401

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	356	DFKDCERRF (SEQ	120.000
		ID NO:55)	
2	334	YFKLSHLQM (SEQ	100.000
		ID NO:248)	
3	180	DPMGQQGSL (SEQ	88.000
		ID NO:59)	
4	163	TPSHHAAQF (SEQ	52.800
		ID NO:228)	
5	327	YPGCNKRYF (SEQ	40.000
		ID NO:250)	
6	285	QYRIHTHGV (SEQ	27.500
		ID NO:175)	
7	424	KFARSDELV (SEQ	25.000
		ID NO:119)	
8	326	AYPGCNKRY (SEQ	25.000
		ID NO:42)	
9	192	QYSVPPPVY (SEQ	25.000
		ID NO:176)	
10	417	RWPSCQKKF (SEQ	22.000
		ID NO:196)	

11	278	TPILCGAQY (SEQ ID NO:227)	12.000
12	10	ALLPAVPSL (SEQ ID NO:34)	11.616
13	141	QPAIRNQGY (SEQ ID NO:170)	11.000
14	303	VPGVAPTLV (SEQ ID NO:242)	11.000
15	219	TPYSSDNLY (SEQ ID NO:231)	10.000
16	39	DFAPPGASA (SEQ ID NO:54)	7.920
17	99	QFTGTAGAC (SEQ ID NO:165)	6.000
18	4	DVRDLNALL (SEQ ID NO:62)	5.760
19	70	SFIKQEPSW (SEQ ID NO:210)	5.500
20	63	PPPPPPHSF (SEQ ID NO:158)	5.280

Table XXXII

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA CW0602

	· · · · · · · · · · · · · · · · · · ·		
			Score (Estimate of Half Time of
1		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	332	KRYFKLSHL (SEQ	9.680
		ID NO:127)	٥
2	239	NQMNLGATL (SEQ	6.600
		ID NO:151)	
3	130	NAPYLPSCL (SEQ ID	6.600
		NO:144)	
4	7	DLNALLPAV (SEQ	6.000
	,	ID NO:58)	
5	441	NMTKLQLAL (SEQ	6.000
		ID NO:149)	
6	225	NLYQMTSQL (SEQ	6.000
		ID NO:147)	

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7	4	DVRDLNALL (SEQ	6.000
		ID NO:62)	
8	3	SDVRDLNAL (SEQ	4.400
		ID NO:206)	
9	10	ALLPAVPSL (SEQ ID	4.000
		_ NO:34)	
10	213	QALLLRTPY (SEQ ID	3.300
		NO:160)	
11	319	EKRPFMCAY (SEQ	3.000
		ID NO:67)	
12	30	GAAQWAPVL (SEQ	2.200
		ID NO:86)	
13	242	NLGATLKGV (SEQ	2.200
		ID NO:146)	
14	292	GVFRGIQDV (SEQ	2.200
		ID NO:103)	
15	207	DSCTGSQAL (SEQ	2.200
	***	ID NO:61)	·
16	362	RRFSRSDQL (SEQ ID	2.200
		NO:187)	
17	439	QRNMTKLQL (SEQ	2.200
		ID NO:173)	
18	295	RGIQDVRRV (SEQ	2.200
		ID NO:179)	
19	423	KKFARSDEL (SEQ	2.200
		ID NO:122)	
20	180	DPMGQQGSL (SEQ	2.200
<u> </u>		ID NO:59)	

Table XXXIII Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Human HLA CW0702

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	319	EKRPFMCAY (SEQ	26.880
		ID NO:67)	
2	326	AYPGCNKRY (SEQ	24.000
		ID NO:42)	

3	40	FAPPGASAY (SEQ ID NO:74)	14.784
4	192	QYSVPPPVY (SEQ ID NO:176)	12.000
5	278	TPILCGAQY (SEQ ID NO:227)	12.000
6	219	TPYSSDNLY (SEQ ID NO:231)	12.000
7	213	QALLLRTPY (SEQ ID NO:160)	8.800
8	125	ARMFPNAPY (SEQ ID NO:38)	8.000
9	327	YPGCNKRYF (SEQ ID NO:250)	6.600
10	152	VTFDGTPSY (SEQ ID NO:244)	5.600
11	141	QPAIRNQGY (SEQ ID NO:170)	4.800
12	345	RKHTGEKPY (SEQ ID NO:184)	4.000
13	185	QGSLGEQQY (SEQ ID NO:166)	4.000
14	101	TGTAGACRY (SEQ ID NO:224)	4.000
15	375	RRHTGVKPF (SEQ ID NO:188)	4.000
16	263	GQSNHSTGY (SEQ ID NO:100)	4.000
17	163	TPSHHAAQF (SEQ ID NO:228)	3.000
18	33	QWAPVLDFA (SEQ ID NO:174)	2.688
19	130	NAPYLPSCL (SEQ ID NO:144)	2.640
20	84	HEEQCLSAF (SEQ ID NO:107)	2.400

Table XXXIV

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Mouse MHC Class I Db

Score (Estimate of Half Time of Subsequence Residue Disassociation of a Molecule Rank **Start Position** Listing Containing This Subsequence) CMTWNQMNL (SEQ 1 235 5255.712 ID NO:49) 2 126 RMFPNAPYL (SEQ 1990.800 ID NO:185) YSSDNLYOM (SEO 930.000 3 221 ID NO:253) 228 OMTSOLECM (SEO 33.701 4 ID NO:169) 5 NQMNLGATL (SEQ 239 21.470 ID NO:151) 6 441 NMTKLQLAL (SEQ 19.908 ID NO:149) 7 437 MHQRNMTKL (SEQ 19.837 ID NO:143) SCLESQPAI (SEQ ID 8 136 11.177 NO:198) HSFKHEDPM (SEQ 9 174 10.800 ID NO:110) RVPGVAPTL (SEQ 10 302 10.088 ID NO:195) 11 130 NAPYLPSCL (SEQ ID 8.400 NO:144) ALLPAVPSL (SEQ ID 12 10 5.988 NO:34) SCTGSQALL (SEQ ID 208 13 4.435 NO:202) 14 209 CTGSQALLL (SEQ ID 3.548 NO:52) WNQMNLGAT (SEQ 15 238 3.300 ID NO:245) RTPYSSDNL (SEQ ID 16 218 3.185 NO:194) 17 24 CALPVSGAA (SEQ 2.851

		ID NO:43)	
18	18	LGGGGGCAL (SEQ ID NO:134).	2.177
19	142	PAIRNQGYS (SEQ ID NO:152)	2.160
20	30	GAAQWAPVL (SEQ ID NO:86)	1.680

Table XXXV

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Mouse MHC Class I Dd

	T		
			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	112	FGPPPPSQA (SEQ ID	48.000
		NO:76)	
2	122	SGQARMFPN (SEQ	36.000
		ID NO:212)	
3	104	AGACRYGPF (SEQ	30.000
		ID NO:31)	
4	218	RTPYSSDNL (SEQ ID	28.800
	1	NO:194)	
5	130	NAPYLPSCL (SEQ ID	20.000
ļ		NO:144)	
6	302	RVPGVAPTL (SEQ	20.000
ŀ		ID NO:195)	
7	18	LGGGGGCAL (SEQ	20.000
		ID NO:134)	
8	81	AEPHEEQCL (SEQ ID	10.000
		NO:30)	
9	29	SGAAQWAPV (SEQ	7.200
		ID NO:211)	
10	423	KKFARSDEL (SEQ	7.200
Ì	1	ID NO:122)	_
11	295	RGIQDVRRV (SEQ	7.200
		ID NO:179)	
12	390	RKFSRSDHL (SEQ ID	6.000
		NO:183)	
13	332	KRYFKLSHL (SEQ	6.000

		ID NO:127)	
14	362	RRFSRSDQL (SEQ ID NO:187)	6.000
15	417	RWPSCQKKF (SEQ ID NO:196)	6.000
16	160	YGHTPSHHA (SEQ ID NO:249)	6.000
17	20	GGGGCALPV (SEQ ID NO:92)	6.000
18	329	GCNKRYFKL (SEQ ID NO:90)	5.000
19	372	RHQRRHTGV (SEQ ID NO:181)	4.500
20	52	GGPAPPPAP (SEQ ID NO:93)	4.000

Table XXXVI

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Mouse MHC Class I Kb

		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	329	GCNKRYFKL (SEQ ID NO:90)	24.000
2	225	NLYQMTSQL (SEQ ID NO:147)	10.000
3	420	SCQKKFARS (SEQ ID NO:200)	3.960
4	218	RTPYSSDNL (SEQ ID NO:194)	3.630
5	437	MHQRNMTKL (SEQ ID NO:143)	3.600
6	387	TCQRKFSRS (SEQ ID NO:219)	3.600
7	302	RVPGVAPTL (SEQ ID NO:195)	3.300
8	130	NAPYLPSCL (SEQ ID NO:144)	3.000
9	289	HTHGVFRGI (SEQ ID	3.000

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		NO:113)	
10	43	PGASAYGSL (SEQ ID NO:153)	2.400
11	155	DGTPSYGHT (SEQ ID NO:56)	2.400
12	273	SDNHTTPIL (SEQ ID NO:204)	2.200
13	126	RMFPNAPYL (SEQ ID NO:185)	2.200
14	128	FPNAPYLPS (SEQ ID NO:79)	2.000
15	3	SDVRDLNAL (SEQ ID NO:206)	1.584
16	207	DSCTGSQAL (SEQ ID NO:61)	1.584
17	332	KRYFKLSHL (SEQ ID NO:127)	1.500
18	18	LGGGGGCAL (SEQ ID NO:134)	1.320
19	233	LECMTWNQM (SEQ ID NO:131)	1.320
20	441	NMTKLQLAL (SEQ ID NO:149)	1.200

Table XXXVII

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Mouse MHC Class I Kd

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	285	QYRIHTHGV (SEQ	600.000
		ID NO:175)	
2	424	KFARSDELV (SEQ	288.000
		ID NO:119)	
3	334	YFKLSHLQM (SEQ	120.000
		ID NO:248)	
4	136	SCLESQPTI (SEQ ID	115.200
		NO:199)	
5	239	NQMNLGATL (SEQ	115.200
		ID NO:151)	
6	10	ALLPAVSSL (SEQ ID	115.200
		NO:35)	
7	47	AYGSLGGPA (SEQ	86.400
		ID NO:41)	
8	180	DPMGQQGSL (SEQ	80.000
		ID NO:59)	
9	270	GYESDNHTA (SEQ	72.000
	ļ	ID NO:105)	
10	326	AYPGCNKRY (SEQ	60.000
		ID NO:42)	
11	192	QYSVPPPVY (SEQ	60.000
		' ID NO:176)	
12	272	ESDNHTAPI (SEQ ID	57.600
		NO:70)	
13	289	HTHGVFRGI (SEQ ID	57.600
		NO:113)	
14	126	DVRDLNALL (SEQ	57.600
		ID NO:62)	
15	4	CTGSQALLL (SEQ ID	57.600
		NO:52)	
16	208	SCTGSQALL (SEQ ID	48.000
		NO:202)	
17	441	NMTKLQLAL (SEQ	48.000
		ID NO:149)	

18	207	DSCTGSQAL (SEQ ID NO:61)	48.000
19	130	NAPYLPSCL (SEQ ID NO:144)	48.000
20	235	CMTWNQMNL (SEQ ID NO:49)	48.000

Table XXXVIII Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Mouse MHC Class I Kk

Score (Estimate of Half Time of Subsequence Residue Disassociation of a Molecule Rank **Start Position** Listing Containing This Subsequence) AEPHEEQCL (SEQ ID 81 40.000 NO:30) EEQCLSAFT (SEQ ID 2 85 40.000 NO:65) 3 429 DELVRHHNM (SEQ 20.000 ID NO:53) SETSEKRPF (SEQ ID 315 20.000 4 NO:209) TEGOSNHST (SEQ ID 20.000 5 261 NO:221) 410 SEKPFSCRW (SEQ 10.000 6 ID NO:207) 7 272 ESDNHTTPI (SEQ ID 10.000 NO:71) 8 318 SEKRPFMCA (SEQ 10.000 ID NO:208) LESQPAIRN (SEQ ID 10.000 9 138 NO:132) 10 233 LECMTWNQM (SEQ 10.000 ID NO:131) 298 **QDVRRVPGV (SEQ** 10.000 11 ID NO:164) HEEQCLSAF (SEQ ID 12 84 10.000 NO:107) **GEKPYQCDF (SEQ** 349 10.000 13 ID NO:91)

14	289	HTHGVFRGI (SEQ ID NO:113)	10.000
15	179	EDPMGQQGS (SEQ ID NO:64)	8.000
16	136	SCLESQPAI (SEQ ID NO:198)	5.000
17	· 280	ILCGAQYRI (SEQ ID NO:116)	5.000
18	273	SDNHTTPIL (SEQ ID NO:204)	4.000
19	428	SDELVRHHN (SEQ ID NO:203)	4.000
20	3	SDVRDLNAL (SEQ ID NO:206)	4.000

Table XXXIX

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Human WT1 Peptides to Mouse MHC Class I Ld

		Subsequence Residue	Score (Estimate of Half Time of Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	163	TPSHHAAQF (SEQ ID NO:228)	360.000
2	327	YPGCNKRYF (SEQ ID NO:250)	300.000
3	180	DPMGQQGSL (SEQ ID NO:59)	150.000
4	26	LPVSGAAQW (SEQ ID NO:138)	93.600
5	278.	TPILCGAQY (SEQ ID NO:227)	72.000
6	141	QPAIRNQGY (SEQ ID NO:170)	60.000
7	219	TPYSSDNLY (SEQ ID NO:231)	60.000
8	303	VPGVAPTLV (SEQ ID NO:242)	60.000
9	120	ASSGQARMF (SEQ ID NO:40)	50.000

10	63	PPPPPHSF (SEQ ID	45.000	
		NO:158)		
11	113	GPPPPSQAS (SEQ ID	45.000	·
		NO:97)		
12	157	TPSYGHTPS (SEQ ID	39.000	
		NO:229)		
13	207	DSCTGSQAL (SEQ	32.500	
		ID NO:61)		
14	110	GPFGPPPPS (SEQ ID	30.000	
		NO:96)		
15	82	EPHEEQCLS (SEQ ID	30.000	
		NO:68)		
16	412	KPFSCRWPS (SEQ ID	30.000	
		NO:123)		
17	418	WPSCQKKFA (SEQ	30.000	
		ID NO:246)		
18	221	YSSDNLYQM (SEQ	30.000	
		ID NO:253)		
19	204	TPTDSCTGS (SEQ ID	30.000	
		NO:230)		
20	128	FPNAPYLPS (SEQ ID	30.000	
		NO:79)		

Table XL

Results of BIMAS HLA Peptide Binding Prediction Analysis for
Binding of Human WT1 Peptides to Cattle HLA A20

Score (Estimate of Half Time of Subsequence Residue Disassociation of a Molecule Containing This Subsequence) **Start Position** Listing Rank 1 350 EKPYQCDFK (SEQ 1000.00 ID NO:66) EKRPFMCAY (SEQ 2 319 500.000 ID NO:67) KKFARSDEL (SEQ 423 500.000 3 ID NO:122) RKHTGEKPY (SEQ 500.000 345 ID NO:184) RKFSRSDHL (SEQ ID 500.000 390 5 NO:183) CLESQPAIR (SEQ ID 137 120.000 6

		NO:47)	
7	380	VKPFQCKTC (SEQ ID NO:239)	100.000
8	407	GKTSEKPFS (SEQ ID NO:95)	100.000
9	335	FKLSHLQMH (SEQ ID NO:78)	100.000
10	247	LKGVAAGSS (SEQ ID NO:135)	100.000
11	370	LKRHQRRHT (SEQ ID NO:136)	100.000
12	258	VKWTEGQSN (SEQ ID NO:240)	100.000
13	398	LKTHTRTHT (SEQ ID NO:137)	100.000
14	331	NKRYFKLSH (SEQ ID NO:145)	100.000
15	357	FKDCERRFS (SEQ ID NO:77)	100.000
16	385	CKTCQRKFS (SEQ ID NO:46)	100.000
17	294	FRGIQDVRR (SEQ ID NO:81)	80.000
18	368	DQLKRHQRR (SEQ ID NO:60)	80.000
19	432	VRHHNMHQR (SEQ ID NO:243)	80.000
20	118	SQASSGQAR (SEQ ID NO:216)	80.000

Table XLI

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Mouse WT1 Peptides to Mouse MHC Class I A 0201

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			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	126	RMFPNAPYL (SEQ	313.968
		ID NO:293)	•
2	187	SLGEQQYSV (SEQ	285.163
L		ID NO:299)	

3	10	ALLPAVSSL (SEQ ID NO:255)	181.794
4	225	NLYQMTSQL (SEQ ID NO:284)	68.360
5	292	GVFRGIQDV (SEQ ID NO:270)	51.790
6	93	TLHFSGQFT (SEQ ID NO:302)	40.986
7	191	QQYSVPPPV (SEQ ID NO:290)	22.566
8	280	ILCGAQYRI (SEQ ID NO:274)	17.736
9	441	NMTKLHVAL (SEQ ID NO:285)	15.428
10	235	CMTWNQMNL (SEQ ID NO:258)	15.428
11	7	DLNALLPAV (SEQ ID NO:261)	11.998
12	242	NLGATLKGM (SEQ ID NO:283)	11.426
13	227	YQMTSQLEC (SEQ ID NO:307)	8.573
14	239	NQMNLGATL (SEQ ID NO:286)	8.014
15	309	TLVRSASET (SEQ ID NO:303)	7.452
16	408	KTSEKPFSC (SEQ ID NO:277)	5.743
17	340	LQMHSRKHT (SEQ ID NO:280)	4.752
18	228	QMTSQLECM (SEQ ID NO:289)	4.044
19	37	VLDFAPPGA (SEQ ID NO:304)	3.378
20	302	RVSGVAPTL (SEQ ID NO:295)	1.869

Table XLII Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Mouse WT1 Peptides to Mouse MHC Class I Db

	·		Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	221	YSSDNLYQM (SEQ	312.000
		ID NO:308)	·
2	126	RMFPNAPYL (SEQ	260.000
		ID NO:293)	
3	235	CMTWNQMNL (SEQ	260.000
		ID NO:258)	
4	437	MHQRNMTKL (SEQ	200.000
		ID NO:281)	
5	238	WNQMNLGAT (SEQ	12.000
		ID NO:305)	
6	130	NAPYLPSCL (SEQ ID	8.580
		NO:282)	·
7	3	SDVRDLNAL (SEQ	7.920
l		ID NO:298)	
8	136	SCLESQPTI (SEQ ID	7.920
	,	NO:296)	
9	81	AEPHEEQCL (SEQ ID	6.600
		NO:254)	
10	10	ALLPAVSSL (SEQ ID	6.600
		NO:255)	
11	218	RTPYSSDNL (SEQ ID	6.000
		NO:294)	
12	441	NMTKLHVAL (SEQ	3.432
		ID NO:285)	
13	228	QMTSQLECM (SEQ	3.120
		ID NO:289)	. 100
14	174	HSFKHEDPM (SEQ	3.120
<u> </u>		ID NO:272)	2 (40
15	242	NLGATLKGM (SEQ	2.640
<u> </u>		ID NO:283)	2.640
16	261	TEGQSNHGI (SEQ ID	2.640
L	<u> </u>	NO:301)	

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17	225	NLYQMTSQL (SEQ ID NO:284)	2.640	
18	207	DSCTGSQAL (SEQ ID NO:263)	2.600	
19	119	QASSGQARM (SEQ ID NO:288)	2.600	:
20	18	LGGGGCGL (SEQ ID NO:279)	2.600	

Table XLIII

Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Mouse WT1 Peptides to Mouse MHC Class I Kb

r——			C (F.4' CH-1CT'. C
			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	329	GCNKRYFKL (SEQ	24.000
		ID NO:268)	
2	225	NLYQMTSQL (SEQ	10.000
[ID NO:284)	
3	420	SCQKKFARS (SEQ	3.960
į		ID NO:297)	
4	218	RTPYSSDNL (SEQ ID	3.630
		NO:294)	
5	437	MHQRNMTKL (SEQ	3.600
	ļ	ID NO:281)	
6	387	TCQRKFSRS (SEQ ID	3.600
		NO:300)	
7	289	HTHGVFRGI (SEQ ID	3.000
		NO:273)	
8	130	NAPYLPSCL (SEQ ID	3.000
		NO:282)	
9	43	PGASAYGSL (SEQ	2.400
1		ID NO:287)	
10	155	DGAPSYGHT (SEQ	2.400
		ID NO:260)	
11	126	RMFPNAPYL (SEQ	2.200
		ID NO:293)	
12	128	FPNAPYLPS (SEQ ID	2.000
		NO:267)	

13	207	DSCTGSQAL (SEQ	1.584
		ID NO:263)	
14	3	SDVRDLNAL (SEQ	1.584
		ID NO:298)	
15	332	KRYFKLSHL (SEQ	1.500
		ID NO:276)	
16	233	LECMTWNQM (SEQ	1.320
		ID NO:278)	
17	18	LGGGGGCGL (SEQ	1.320
		ID NO:279)	
18	242	NLGATLKGM (SEQ	1.200
		ID NO:283)	
19	123	GQARMFPN (SEQ ID	1.200
		NO:269)A	
20	441	NMTKLHVAL (SEQ	1.200
		ID NO:285)	

Table XLIV Results of BIMAS HLA Peptide Binding Prediction Analysis for Binding of Mouse WT1 Peptides to Mouse MHC Class I Kd

			Score (Estimate of Half Time of
		Subsequence Residue	Disassociation of a Molecule
Rank	Start Position	Listing	Containing This Subsequence)
1	285	QYRIHTHGV (SEQ	600.000
		ID NO:291)	
2	424	KFARSDELV (SEQ	288.000
		ID NO:275)	
3	334	YFKLSHLQM (SEQ	120.000
		ID NO:306)	
4	136	SCLESQPTI (SEQ ID	115.200
		NO:296)	
5	239	NQMNLGATL (SEQ	115.200
		ID NO:286)	
6	10	ALLPAVSSL (SEQ ID	115.200
		NO:255)	
7	47	AYGSLGGPA (SEQ	86.400
		ID NO:256)	
8	180	DPMGQQGSL (SEQ	80.000
		ID NO:262)	

9	270	GYESDNHTA (SEQ	GYESDNHTA (SEQ 72.000 ID NO:271)	
 10 			40.000	
10	192	QYSVPPPVY (SEQ	60.000	
		ID NO:292)		
11	326	AYPGCNKRY (SEQ	60.000	
		ID NO:257)		
12	289	HTHGVFRGI (SEQ ID	57.600	
		NO:273)		
13	4	DVRDLNALL (SEQ	57.600	
		ID NO:264)		
14	126	RMFPNAPYL (SEQ	57.600	
		ID NO:293)		
15	209	CTGSQALLL (SEQ ID	48.000	
		NO:259)		
16	86	EQCLSAFTL (SEQ ID	48.000	
,		NO:265)		
17	302	RVSGVAPTL (SEQ	48.000	
		ID NO:295)		
18	218	RTPYSSDNL (SEQ ID	48.000	
	-	NO:294)		
19	272	ESDNHTAPI (SEQ ID	48.000	
	_ · _	NO:266)		
20	225	NLYQMTSQL (SEQ	48.000	
~	223	ID NO:284)	,0.000	
		10 110.207)		

Table XLV Results of TSites Peptide Binding Prediction Analysis for Human WT1 Peptides Capable of Eliciting a Helper T cell Response

Peptide	Sequence
p6-23	RDLNALLPAVPSLGGGG (SEQ ID NO:1)
p30-35	GAAQWA (SEQ ID NO:309)
p45-56	ASAYGSLGGPAP (SEQ ID NO:310)
p91-105	AFTVHFSGQFTGTAG (SEQ ID NO:311)
p117-139	PSQASSGQARMFPNAPYLPSCLE (SEQ ID NO:2)
p167-171	HAAQF (SEQ ID NO:312)
p202-233	CHTPTDSCTGSQALLLRTPYSSDNLYQMTSQL (SEQ ID NO:313)
p244-262	GATLKGVAAGSSSSVKWTE (SEQ ID NO:4)
p287-318	RIHTHGVFRGIQDVRRVPGVAPTLVRSASETS (SEQ ID NO:314)

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p333-336	RYFK (SEQ ID NO:315)
p361-374	ERRFSRSDQLKRHQ (SEQ ID NO:316)
p389-410	QRKFSRSDHLKTHTRTHTGKTS (SEQ ID NO:317)
p421-441	CQKKFARSDELVRHHNMHQRN (SEQ ID NO:318)

Certain CTL peptides (shown in Table XLVI) were selected for further study. For each peptide in Table XLVI, scores obtained using BIMAS HLA peptide binding prediction analysis are provided.

Table XLVI
WT1 Peptide Sequences and HLA Peptide Binding Predictions

Peptide	Sequence	Comments
p329-337	GCNKRYFKL	Score 24,000
	(SEQ ID NOs: 90 and	
	268)	
p225-233	NLYQMTSQL	binds also to class II and HLA A2, Kd,
	(SEQ ID NOs: 147 and	score 10,000
	284)	·
p235-243	CMTWNQMNL	binds also to HLA A2, score 5,255,712
	(SEQ ID NOs: 49 and	·
·	258)	
p126-134	RMFPNAPYL	binds also to Kd, class II and HLA A2,
	(SEQ ID NOs: 185 and	score 1,990,800
	293)	
p221-229	YSSDNLYQM	binds also to Ld, score 312,000
	(SEQ ID NOs: 253 and	
	308)	
p228-236	QMTSQLECM	score 3,120
	(SEQ ID NOs: 169 and	
	289)	
p239-247	NQMNLGATL	binds also to HLA A 0201, Kd, score
	(SEQ ID NOs: 151 and	8,015
	286)	
mouse p136-144	SCLESQPTI	binds also to Kd, 1 mismatch to human
	(SEQ ID NO:296)	
human p136-144	SCLESQPAI	score 7,920
	(SEQ ID NO:198)	
mouse p10-18	ALLPAVSSL	binds also to Kd, HLA A2, 1 mismatch

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	(SEQ ID NO:255)	to human
human p10-18	ALLPAVPSL	score 6,600
	(SEQ ID NO:34)	

Peptide binding to C57Bl/6 murine MHC was confirmed using the leukemia cell line RMA-S, as described by Ljunggren et al., *Nature 346*:476-480, 1990. In brief, RMA-S cells were cultured for 7 hours at 26°C in complete medium supplemented with 1% FCS. A total of 10⁶ RMA-S cells were added into each well of a 24-well plate and incubated either alone or with the designated peptide (25ug/ml) for 16 hours at 26°C and additional 3 hours at 37°C in complete medium. Cells were then washed three times and stained with fluorescein isothiocyanate-conjugated anti D^b or anti-K^b antibody (PharMingen, San Diego, CA). Labeled cells were washed twice, resuspended and fixed in 500ul of PBS with 1% paraformaldehyde and analyzed for fluorescence intensity in a flow cytometer (Becton-Dickinson FACSCalibur®). The percentage of increase of D^b or K^b molecules on the surface of the RMA-S cells was measured by increased mean fluorescent intensity of cells incubated with peptide compared with that of cells incubated in medium alone.

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Mice were immunized with the peptides capable of binding to murine class I MHC. Following immunization, spleen cells were stimulated *in vitro* and tested for the ability to lyse targets incubated with WT1 peptides. CTL were evaluated with a standard chromium release assay (Chen et al., *Cancer Res. 54*:1065-1070, 1994). 10⁶ target cells were incubated at 37°C with 150μCi of sodium ⁵¹Cr for 90 minutes, in the presence or absence of specific peptides. Cells were washed three times and resuspended in RPMI with 5% fetal bovine serum. For the assay, 10⁴ ⁵¹Cr-labeled target cells were incubated with different concentrations of effector cells in a final volume of 200μl in U-bottomed 96-well plates. Supernatants were removed after 4 to 7 hours at 37°C, and the percentage specific lysis was determined by the formula:

% specific lysis = 100 x (experimental release - spontaneous release)/(maximum release-spontaneous release).

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The results, presented in Table XLVII, show that some WT1 peptides can bind to class I MHC molecules, which is essential for generating CTL. Moreover, several of the peptides were able to elicit peptide specific CTL (Figures 9A and 9B), as determined using chromium release assays. Following immunization to CTL peptides p10-18 human, p136-144 human, p136-144 mouse and p235-243, peptide specific CTL lines were generated and clones were established. These results indicate that peptide specific CTL can kill malignant cells expressing WT1.

<u>Table XLVII</u>

<u>Binding of WT1 CTL Peptides to mouse B6 class I antigens</u>

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Peptide	Binding Affinity to Mouse MHC Class I
Positive control	91%
negative control	0.51.3%
p235-243	33.6%
p136-144 mouse	27.9%
p136-144 human	52%
p10-18: human	2.2%
p225-233	5.8%
p329-337	1.2%
p126-134	0.9%
p221-229	0.8%
p228-236	1.2%
p239-247	1%

Example 5 Use of a WT1 Polypeptide to Elicit WT1 Specific CTL in Mice

This Example illustrates the ability of a representative WT1 polypeptide to elicit CTL immunity capable of killing WT1 positive tumor cell lines.

P117-139, a peptide with motifs appropriate for binding to class I and class II MHC, was identified as described above using TSITES and BIMAS HLA peptide binding prediction analyses. Mice were immunized as described in Example 3. Following

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immunization, spleen cells were stimulated *in vitro* and tested for the ability to lyse targets incubated with WT1 peptides, as well as WT1 positive and negative tumor cells. CTL were evaluated with a standard chromium release assay. The results, presented in Figures 10A-10D, show that P117 can elicit WT1 specific CTL capable of killing WT1 positive tumor cells, whereas no killing of WT1 negative cells was observed. These results demonstrate that peptide specific CTL in fact kill malignant cells expressing WT1 and that vaccine and T cell therapy are effective against malignancies that express WT1.

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Similar immunizations were performed using the 9-mer class I MHC binding peptides p136-144, p225-233, p235-243 as well as the 23-mer peptide p117-139. Following immunization, spleen cells were stimulated *in vitro* with each of the 4 peptides and tested for ability to lyse targets incubated with WT1 peptides. CTL were generated specific for p136-144, p235-243 and p117-139, but not for p225-233. CTL data for p235-243 and p117-139 are presented in Figures 11A and 11B. Data for peptides p136-144 and p225-233 are not depicted.

CTL lysis demands that the target WT1 peptides are endogenously processed and presented in association with tumor cell class I MHC molecules. The above WT1 peptide specific CTL were tested for ability to lyse WT1 positive versus negative tumor cell lines. CTL specific for p235-243 lysed targets incubated with the p235-243 peptides, but failed to lyse cell lines that expressed WT1 proteins (Figure 11A). By marked contrast, CTL specific for p117-139 lysed targets incubated with p117-139 peptides and also lysed malignant cells expressing WT1 (Figure 11B). As a negative control, CTL specific for p117-139 did not lyse WT1 negative EL-4 (also referred to herein as E10).

Specificity of WT1 specific lysis was confirmed by cold target inhibition (Figures 12A-12B). Effector cells were plated for various effector: target ratios in 96-well U-bottom plates. A ten-fold excess (compared to hot target) of the indicated peptide-coated target without ⁵¹Cr labeling was added. Finally, 10⁴ ⁵¹Cr-labeled target cells per well were added and the plates incubated at 37°C for 4 hours. The total volume per well was 200µl.

Lysis of TRAMP-C by p117-139 specific CTL was blocked from 58% to 36% by EL-4 incubated with the relevant peptide p117-139, but not with EL-4 incubated with an irrelevant peptide (Figure 12A). Similarly, lysis of BLK-SV40 was blocked from 18% to 0% by EL-4 incubated with the relevant peptide p117-139 (Figure 12B). Results validate that WT1 peptide specific CTL specifically kill malignant cells by recognition of processed WT1.

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Several segments with putative CTL motifs are contained within p117-139. To determine the precise sequence of the CTL epitope all potential 9-mer peptides within p117-139 were synthesized (Table XLVIII). Two of these peptides (p126-134 and p130-138) were shown to bind to H-2^b class I molecules (Table XLVIII). CTL generated by immunization with p117-139 lysed targets incubated with p126-134 and p130-138, but not the other 9-mer peptides within p117-139 (Figure 13A).

The p117-139 specific CTL line was restimulated with either p126-134 or p130-138. Following restimulation with p126-134 or p130-138, both T cell lines demonstrated peptide specific lysis, but only p130-138 specific CTL showed lysis of a WT1 positive tumor cell line (Figures 13B and 13C). Thus, p130-138 appears to be the naturally processed epitope.

Table XLVIII

Binding of WT1 CTL 9mer Peptides within p117-139 to mouse B6 class I antigens

Peptide				Binding Affinity to Mouse MHC Class I
P117-125	PSQASSGQA	(SEQ	ID	2%
NO:221)				
P118-126	SQASSGQAR	(SEQ	ID	2%
NO:216)				
P119-127	QASSGQARM	(SEQ	ID	2%
NOs: 161 and	d 288)			
P120-128	ASSGQARMF	(SEQ	ID	1%
NO:40				
P121-129	SSGQARMFP	(SEQ	ID	1%
NO:222)		_		
P122-130	SGQARMFPN	(SEQ	ID	1%

NO:212)			
P123-131	GQARMFPNA (SEQ	ID	1%
NOs: 98 and 2	269)		
P124-132	QARMFPNAP (SEQ	ID	1% ·
NO:223)			
P125-133	ARMFPNAPY (SEQ	ID	1%
NO:38)			
P126-134	RMFPNAPYL (SEQ	ID	79%
NOs: 185 and	1 293)		
P127-135	MFPNAPYLP (SEQ	ID	2%
NO:224)			
P128-136	FPNAPYLPS (SEQ	ID	1%
NOs: 79 and	267)		
P129-137	PNAPYLPSC (SEQ	ID	1%
NO:225)			
P130-138	NAPYLPSCL (SEQ	ID	79%
NOs: 144 and	1 282)		
P131-139	APYLPSCLE (SEQ	ID	1%
NO:226)			

Example 6 Identification of WT1 Specific mRNA in Mouse Tumor Cell Lines

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This Example illustrates the use of RT-PCR to detect WT1 specific mRNA in cells and cell lines.

Mononuclear cells were isolated by density gradient centrifugation, and were immediately frozen and stored at -80°C until analyzed by RT-PCR for the presence of WT1 specific mRNA. RT-PCR was generally performed as described by Fraizer et al., Blood 86:4704-4706, 1995. Total RNA was extracted from 10⁷ cells according to standard procedures. RNA pellets were resuspended in 25 μL diethylpyrocarbonate treated water and used directly for reverse transcription. The zinc-finger region (exons 7 to 10) was amplified by PCR as a 330 bp mouse cDNA. Amplification was performed in a thermocycler during one or, when necessary, two sequential rounds of PCR. AmpliTaq

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DNA Polymerase (Perkin Elmer Cetus, Norwalk, CT), 2.5 mM MgCl₂ and 20 pmol of each primer in a total reaction volume of 50µl were used. Twenty µL aliquots of the PCR products were electrophoresed on 2% agarose gels stained with ethidium bromide. The gels were photographed with Polaroid film (Polaroid 667, Polaroid Ltd., Hertfordshire, England). Precautions against cross contamination were taken following the recommendations of Kwok and Higuchi, *Nature 339*:237-238, 1989. Negative controls included the cDNA- and PCR-reagent mixes with water instead of cDNA in each experiment. To avoid false negatives, the presence of intact RNA and adequate cDNA generation was evaluated for each sample by a control PCR using β-actin primers. Samples that did not amplify with these primers were excluded from analysis.

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Primers for amplification of WT1 in mouse cell lines were: P115: 1458-1478: 5' CCC AGG CTG CAA TAA GAG ATA 3' (forward primer; SEQ ID NO:21); and P116: 1767-1787: 5' ATG TTG TGA TGG CGG ACC AAT 3' (reverse primer; SEQ ID NO:22) (see Inoue et al, Blood 88:2267-2278, 1996; Fraizer et al., Blood 86:4704-4706, 1995).

Beta Actin primers used in the control reactions were: 5' GTG GGG CGC CCC AGG CAC CA 3' (sense primer; SEQ ID NO:23); and 5' GTC CTT AAT GTC ACG CAC GAT TTC 3' (antisense primer; SEQ ID NO:24)

Primers for use in amplifying human WT1 include: P117: 954-974: 5' GGC ATC TGA GAC CAG TGA GAA 3' (SEQ ID NO:25); and P118: 1434-1414: 5' GAG AGT CAG ACT TGA AAG CAGT 3' (SEQ ID NO:5). For nested RT-PCR, primers may be: P119: 1023-1043: 5' GCT GTC CCA CTT ACA GAT GCA 3' (SEQ ID NO:26); and P120: 1345-1365: 5' TCA AAG CGC CAG CTG GAG TTT 3' (SEQ ID NO:27).

Table XLVIII shows the results of WT1 PCR analysis of mouse tumor cell lines. Within Table IV, (+++) indicates a strong WT1 PCR amplification product in the first step RT PCR, (++) indicates a WT1 amplification product that is detectable by first step WT1 RT PCR, (+) indicates a product that is detectable only in the second step of WT1 RT PCR, and (-) indicates WT1 PCR negative.

<u>Table XLIX</u> <u>Detection of WT1 mRNA in Mouse Tumor Cell Lines</u>

Cell Line	WT1 mRNA
K562 (human leukemia; ATCC): Positive control; (Lozzio and Lozzio, <i>Blood 45</i> :321-334, 1975)	11+
TRAMPC (SV40 transformed prostate, B6); Foster et al., Cancer Res. 57:3325-3330, 1997	+++
BLK-SV40 HD2 (SV40-transf. fibroblast, B6; ATCC); <i>Nature</i> 276:510-511, 1978	++
CTLL (T-cell, B6; ATCC); Gillis, Nature 268:154-156, 1977)	+
FM (FBL-3 subline, leukemia, B6); Glynn and Fefer, Cancer Res. 28:434-439, 1968	+
BALB 3T3 (ATCC); Aaroston and Todaro, J. Cell. Physiol. 72:141-148, 1968	+
S49.1 (Lymphoma, T-cell like, B/C; ATCC); Horibata and	+
Harris, Exp. Cell. Res. 60:61, 1970	
BNL CL.2 (embryonic liver, B/C; ATCC); Nature 276:510-511,	+
1978	
MethA (sarcoma, B/C); Old et al., Ann. NY Acad. Sci. 101:80-106, 1962	-
P3.6.2.8.1 (myeloma, B/C; ATCC); Proc. Natl. Acad. Sci. USA 66:344, 1970	-
P2N (leukemia, DBA/2; ATCC); Melling et al., <i>J. Immunol.</i> 117:1267-1274, 1976	_
BCL1 (lymphoma, B/C; ATCC); Slavin and Strober, Nature 272:624-626, 1977	- 1
LSTRA (lymphoma, B/C); Glynn et al., Cancer Res. 28:434-439, 1968	-
E10/EL-4 (lymphoma, B6); Glynn et al., Cancer Res. 28:434-439, 1968	_

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From the foregoing it will be appreciated that, although specific embodiments of the invention have been described herein for purposes of illustration, various modifications may be made without deviating from the spirit and scope of the invention. Accordingly, the invention is not limited except as by the appended claims.

CLAIMS

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- A polypeptide comprising an immunogenic portion of a native 1. WT1, or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with WT1-specific antisera and/or T-cell lines or clones is not substantially diminished, wherein the polypeptide comprises no more than 16 consecutive amino acid residues present within a native WT1 polypeptide.
- 2. A polypeptide according to claim 1, wherein the immunogenic portion binds to an MHC class I molecule.
- A polypeptide according to claim 1, wherein the immunogenic 3. portion binds to an MHC class II molecule.
- A polypeptide according to claim 1, wherein the polypeptide 4. comprises a sequence selected from the group consisting of:
 - sequences recited in one or more of Tables II XLVI; (a)
- variants of the foregoing sequences that differ in one or more (b) substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera and/or T-cell lines or clones is not substantially diminished; and
- mimetics of the foregoing sequences, wherein the ability of the mimetic to react with antigen-specific antisera and/or T-cell lines or clones is not substantially diminished.
- A polypeptide according to claim 1, wherein the polypeptide 5. comprises a sequence selected from the group consisting of:
- ALLPAVPSL (SEQ ID NO:34), GATLKGVAA (SEQ ID (a) NO:88), CMTWNQMNL (SEQ ID NOs: 49 and 258), SCLESQPTI (SEQ ID NOs: 199 and 296), SCLESQPAI (SEQ ID NO:198), NLYQMTSQL (SEQ ID NOs: 147 and

284); ALLPAVSSL (SEQ ID NOs: 35 and 255), RMFPNAPYL (SEQ ID NOs: 185 and 293);

- (b) variants of the foregoing sequences that differ in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera and/or T-cell lines or clones is not substantially diminished; and
- (c) mimetics of the foregoing sequences, wherein the ability of the mimetic to react with antigen-specific antisera and/or T-cell lines or clones is not substantially diminished.
- 6. A polypeptide according to claim 1, wherein the polypeptide comprises 4-16 consecutive amino acids of a native WT1 polypeptide.
- 7. A polypeptide according to claim 1, wherein the polypeptide comprises 8-10 consecutive amino acids of a native WT1 polypeptide.
- 8. A polypeptide comprising an immunogenic portion of amino acid residues 1 174 of a native WT1 polypeptide, or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with WT1-specific T-cell lines or clones is not substantially diminished, wherein the polypeptide comprises no more than 16 consecutive amino acid residues present within amino acids 175 to 449 of the native WT1 polypeptide.
- 9. A polypeptide comprising a variant of an immunogenic portion of WT1 that differs in substitutions at between 1 and 3 amino acid positions within the immunogenic portion, such that the ability of the variant to react with WT1-specific antisera and/or T-cell lines or clones is enhanced relative to a native WT1.
- wherein at least one amino acid residue is replaced by a compound that is not an amino

acid, such that the ability of the mimetic to react with antigen-specific antisera and/or T-cell lines or clones is not diminished.

- 11. A pharmaceutical composition comprising a polypeptide according to claim 1, in combination with a pharmaceutically acceptable carrier or excipient.
- 12. A pharmaceutical composition according to claim 11, wherein the polypeptide comprises 4-16 consecutive amino acids of a native WT1 polypeptide.
- 13. A pharmaceutical composition according to claim 11, wherein the polypeptide comprises 8-16 consecutive amino acids of a native WT1 polypeptide.
- 14. A pharmaceutical composition comprising a polypeptide according to claim 8, in combination with a pharmaceutically acceptable carrier or excipient.
- 15. A vaccine comprising a polypeptide according to claim 1, in combination with a non-specific immune response enhancer.
- 16. A vaccine according to claim 15, wherein the polypeptide comprises 4-16 consecutive amino acids of a native WT1 polypeptide.
- 17. A vaccine according to claim 15, wherein the polypeptide comprises 8-10 consecutive amino acids of a native WT1 polypeptide.
- 18. A vaccine according to claim 15, wherein the immune response enhancer is an adjuvant.
- 19. A vaccine comprising a polypeptide according to claim 8, in combination with a non-specific immune response enhancer.

- 20. A vaccine according to claim 19, wherein the immune response enhancer is an adjuvant.
 - 21. A vaccine comprising:
- (a) a WT1 polypeptide, wherein the polypeptide comprises an immunogenic portion of a native WT1 or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific T cell lines or clones is not substantially diminished; and
- (b) a non-specific immune response enhancer that preferentially enhances a T cell response in a patient.
- 22. A vaccine according to claim 21, wherein the immune response enhancer is selected from the group consisting of Montanide ISA50, Seppic MONTANIDE ISA 720, cytokines (e.g., GM-CSF, Flat3-ligand), microspheres, dimethyl dioctadecyl ammoniumbromide (DDA) based adjuvants, AS-1, AS-2, Ribi Adjuvant system based adjuvants, QS21, saponin based adjuvants, Syntex adjuvant in its microfluidized form, MV, ddMV, immune stimulating complex (iscom) based adjuvants and inactivated toxins.
- 23. A pharmaceutical composition comprising a mimetic according to claim 10, in combination with a pharmaceutically acceptable carrier or excipient.
- 24. A vaccine comprising a mimetic according to claim 10, in combination with a non-specific immune response enhancer.
- 25. A polynucleotide encoding a polypeptide according to claim 1 or claim 8.
 - 26. A pharmaceutical composition, comprising:

- (a) a polynucleotide encoding a WT1 polypeptide, wherein the polypeptide comprises an immunogenic portion of a native WT1 or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antibodies and/or T cell lines or clones is not substantially diminished; and
 - (b) a pharmaceutically acceptable carrier or excipient.
 - 27. A pharmaceutical composition, comprising:
- (a) an antibody or antigen-binding fragment thereof that specifically binds to a WT1 polypeptide; and
 - (b) a pharmaceutically acceptable carrier or excipient.
 - 28. A pharmaceutical composition, comprising:
 - (a) a T cell that specifically reacts with a WT1 polypeptide; and
 - (b) a pharmaceutically acceptable carrier or excipient.
 - 29. A pharmaceutical composition, comprising:
 - (a) an antigen presenting cell that expresses
- (i) a WT1 polypeptide that comprises an immunogenic portion of a native WT1 or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antibodies and/or T cell lines or clones is not substantially diminished; and
 - (b) a pharmaceutically acceptable carrier or excipient.
 - 30. A vaccine, comprising:
- (a) a polynucleotide encoding a WT1 polypeptide, wherein the polypeptide comprises an immunogenic portion of a native WT1 or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antibodies and/or T cell lines or clones is not substantially diminished; and

- (b) a non-specific immune response enhancer.
- 31. A vaccine, comprising:
- (a) an antigen presenting cell that expresses:
- (i) a WT1 polypeptide that comprises an immunogenic portion of a native WT1 or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antibodies and/or T cell lines or clones is not substantially diminished; and
 - (b) a non-specific immune response enhancer.
 - 32. A vaccine comprising:
- (a) an anti-idiotypic antibody or antigen-binding fragment thereof that is specifically bound by an antibody that specifically binds to an immunogenic portion of WT1; and
 - (b) non-specific immune response enhancer.
- 33. A vaccine according to any one of claims 30-32, wherein the immune response enhancer is an adjuvant.
- 34. A vaccine according to any one of claims 30-32, wherein the immune response enhancer preferentially enhances a T cell response in a patient.
- 35. A method for enhancing or inducing an immune response in a human patient, comprising administering to a patient a pharmaceutical composition comprising:
- (a) a WT1 polypeptide that comprises an immunogenic portion of a native WT1 or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigenspecific antibodies and/or T cell lines or clones is not substantially diminished; and
 - (b) a physiologically acceptable carrier or excipient;

and thereby enhancing or inducing an immune response specific for WT1 or a cell expressing WT1 in the human patient.

- 36. A method for enhancing or inducing an immune response in a patient, comprising administering to a patient a pharmaceutical composition according to any one of claims 11, 14, 23 or 26-29.
- 37. A method for enhancing or inducing an immune response in a human patient, comprising administering to a patient a vaccine comprising:
- (a) a WT1 polypeptide that comprises an immunogenic portion of a native WT1 or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigenspecific antibodies and/or T cell lines or clones is not substantially diminished; and
- (b) a non-specific immune response enhancer;and thereby enhancing or inducing an immune response specific forWT1 or a cell expressing WT1 in the human patient.
- 38. A method for enhancing or inducing an immune response in a patient, comprising administering to a patient a vaccine according to any one of claims 15, 19, 21, 24 or 30-32, and thereby enhancing or inducing an immune response specific for WT1 or a cell expressing WT1 in the patient.
- 39. A method for inhibiting the development of a malignant disease associated with WT1 expression in a human patient, comprising administering to a human patient a pharmaceutical composition comprising:
- (a) a WT1 polypeptide that comprises an immunogenic portion of a native WT1 or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigenspecific antibodies and/or T cell lines or clones is not substantially diminished; and
 - (b) a physiologically acceptable carrier or excipient;

and thereby inhibiting the development of a malignant disease associated with WT1 expression in the human patient.

- 40. A method for inhibiting the development of a malignant disease associated with WT1 expression in a patient, comprising administering to a patient a pharmaceutical composition according to any one of claims 11, 14, 23 or 26-29, and thereby inhibiting the development of a malignant disease in the patient.
- 41. A method for inhibiting the development of a malignant disease associated with WT1 expression in a human patient, comprising administering to a patient a vaccine comprising:
- (a) a WT1 polypeptide that comprises an immunogenic portion of a native WT1 or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigenspecific antibodies and/or T cell lines or clones is not substantially diminished; and
- (b) a non-specific immune response enhancer; and thereby inhibiting the development of a malignant disease in the patient.
- 42. A method for inhibiting the development of a malignant disease associated with WT1 expression in a patient, comprising administering to a patient a vaccine according to any one of claims 15, 19, 21, 24 or 30-32, and thereby inhibiting the development of a malignant disease in the patient.
- 43. A method according to claim 39 or claim 41, wherein the malignant disease is a leukemia.
- 44. A method according to claim 43, wherein the leukemia is acute myeloid leukemia, acute lymphocytic leukemia or chronic myeloid leukemia.

malignant disease is a cancer.

45. A method according to claim 39 or claim 41, wherein the

- 46. A method according to claim 45, wherein the cancer is breast, lung, thyroid or gastrointestinal cancer or a melanoma.
- 47. A method according to claim 40, wherein the malignant disease is a leukemia.
- 48. A method according to claim 47, wherein the leukemia is acute myeloid leukemia, acute lymphocytic leukemia or chronic myeloid leukemia.
- 49. A method according to claim 40, wherein the malignant disease is a cancer.
- 50. A method according to claim 49, wherein the cancer is breast, lung, thyroid or gastrointestinal cancer or a melanoma.
- 51. A method according to claim 42, wherein the malignant disease is a leukemia.
- 52. A method according to claim 51, wherein the leukemia is acute myeloid leukemia, acute lymphocytic leukemia or chronic myeloid leukemia.
- 53. A method according to claim 42, wherein the malignant disease is a cancer.
- 54. A method according to claim 53, wherein the cancer is breast, lung, thyroid or gastrointestinal cancer or a melanoma.
- 55. A method according to claim 39, wherein the pharmaceutical composition comprises a WT1 polypeptide that comprises a sequence selected from the

group consisting of sequences recited in one or more of Tables II - XLVI and variants of the foregoing sequences that differ in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera and/or T-cell lines or clones is not diminished.

- 56. A method according to claim 39, wherein the pharmaceutical composition comprises a WT1 polypeptide that comprises a sequence selected from the group consisting of ALLPAVPSL (SEQ ID NO:34), GATLKGVAA (SEQ ID NO:88), CMTWNQMNL (SEQ ID NOs: 49 and 258), SCLESQPTI (SEQ ID NOs: 199 and 296), SCLESQPAI (SEQ ID NO:198), NLYQMTSQL (SEQ ID NOs: 147 and 284), ALLPAVSSL (SEQ ID NOs: 35 and 255); RMFPNAPYL (SEQ ID NOs: 185 and 293) and variants of the foregoing sequences that differ in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera and/or T-cell lines or clones is not diminished.
- 57. A method according to claim 41, wherein the vaccine comprises a WT1 polypeptide that comprises a sequence selected from the group consisting of sequences recited in one or more of Tables II XLVI and variants of the foregoing sequences that differ in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera and/or T-cell lines or clones is not diminished.
- 58. A method according to claim 41, wherein the vaccine comprises a WT1 polypeptide that comprises a sequence selected from the group consisting of ALLPAVPSL (SEQ ID NO:34), GATLKGVAA (SEQ ID NO:88), CMTWNQMNL (SEQ ID NOs: 49 and 258), SCLESQPTI (SEQ ID NOs: 199 and 296), SCLESQPAI (SEQ ID NO:198), NLYQMTSQL (SEQ ID NOs: 147 and 284), ALLPAVSSL (SEQ ID NOs: 35 and 255), RMFPNAPYL (SEQ ID NOs: 185 and 293) and variants of the foregoing sequences that differ in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera and/or T-cell lines or clones is not diminished.

- 59. A method for removing cells expressing WT1 from bone marrow, peripheral blood, or a fraction of bone marrow or peripheral blood, comprising contacting bone marrow, peripheral blood or a fraction of bone marrow or peripheral blood with T cells that specifically react with a WT1 polypeptide, wherein the step of contacting is performed under conditions and for a time sufficient to permit the removal of WT1 positive cells to less than 10% of the number of myeloid or lymphatic cells in the bone marrow, peripheral blood or a fraction of bone marrow or peripheral blood.
- 60. A method for inhibiting the development of a malignant disease associated with WT1 expression in a patient, comprising administering to a patient bone marrow, peripheral blood or a fraction or bone marrow or peripheral blood prepared according to the method of claim 59.
- 61. A method according to claim 60, wherein the bone marrow, peripheral blood or fraction is autologous.
- 62. A method according to claim 60, wherein the bone marrow, peripheral blood or fraction is syngeneic or allogeneic.
- 63. A method for stimulating and/or expanding T cells, comprising contacting T cells with a WT1 polypeptide, a polynucleotide encoding a WT1 polypeptide and/or an antigen presenting cell that expresses a WT1 polypeptide under conditions and for a time sufficient to permit the stimulation and/or expansion of T cells.
- 64. A method according to claim 63, wherein the T cells are present within bone marrow, peripheral blood or a fraction of bone marrow or peripheral blood.

- 65. A method according to claim 63, wherein the bone marrow, peripheral blood or fraction is obtained from a patient afflicted with a malignant disease associated with WT1 expression.
- 66. A method according to claim 63, wherein the bone marrow, peripheral blood or fraction is obtained from a mammal that is not afflicted with a malignant disease associated with WT1 expression.
- 67. A method according to claim 63, wherein the T cells are cloned prior to expansion.
- 68. A method for stimulating and/or expanding T cells in a mammal, comprising administering to a mammal a pharmaceutical composition comprising:
 - (a) one or more of:
 - (i) a WT1 polypeptide;
 - (ii) a polynucleotide encoding a WT1 polypeptide; or
- (iii) an antigen-presenting cell that expresses a WT1 polypeptide; and
 - (b) a physiologically acceptable carrier or excipient; and thereby stimulating and/or expanding T cells in a mammal.
- 69. A method for stimulating and/or expanding T cells in a mammal, comprising administering to a mammal a vaccine comprising:
 - (a) one or more of:
 - (i) a WT1 polypeptide;
 - (ii) a polynucleotide encoding a WT1 polypeptide; or
- (iii) an antigen-presenting cell that expresses a WT1 polypeptide; and
 - (b) a non-specific immune response enhancer; and thereby stimulating and/or expanding T cells in a mammal.

- 70. A method for inhibiting the development of a malignant disease associated with WT1 expression in a patient, comprising administering to a patient T cells prepared according to the method of claim 63.
- 71. A method according to claim 70, wherein the bone marrow, peripheral blood or fraction is obtained from a patient afflicted with a malignant disease associated with WT1 expression.
- 72. A method according to claim 70, wherein the bone marrow, peripheral blood or fraction is obtained from a mammal that is not afflicted with a malignant disease associated with WT1 expression.
- 73. A method for monitoring the effectiveness of an immunization or therapy for a malignant disease associated with WT1 expression in a patient, comprising the steps of:
 - (a) incubating a first biological sample with one or more of:
 - (i) a WT1 polypeptide;
 - (ii) a polynucleotide encoding a WT1 polypeptide; or
- (iii) an antigen-presenting cell that expresses a WT1 polypeptide

wherein the first biological sample is obtained from a patient prior to a therapy or immunization, and wherein the incubation is performed under conditions and for a time sufficient to allow immunocomplexes to form;

- (b) detecting immunocomplexes formed between the WT1 polypeptide and antibodies in the biological sample that specifically bind to the WT1 polypeptide;
- (c) repeating steps (a) and (b) using a second biological sample obtained from the patient following therapy or immunization; and

- (d) comparing the number of immunocomplexes detected in the first and second biological samples, and therefrom monitoring the effectiveness of the therapy or immunization in the patient.
- 74. A method according to claim 73, wherein the step of detecting comprises (a) incubating the immunocomplexes with a detection reagent that is capable of binding to the immunocomplexes, wherein the detection reagent comprises a reporter group, (b) removing unbound detection reagent, and (c) detecting the presence or absence of the reporter group.
- 75. A method according to claim 74, wherein the detection reagent comprises a second antibody, or antigen-binding fragment thereof, capable of binding to the antibodies that specifically bind to the WT1 polypeptide.
- 76. A method according to claim 74, wherein the detection reagent comprises Protein A.
- 77. A method according to claim 74, wherein the reporter group is selected from the group consisting of radioisotopes, fluorescent groups, luminescent groups, enzymes, biotin and dye particles.
- 78. A method according to claim 73 wherein a reporter group is bound to the WT1 polypeptide, and wherein the step of detecting comprises removing unbound WT1 polypeptide and subsequently detecting the presence or absence of the reporter group.
- 79. A method for monitoring the effectiveness of an immunization or therapy for a malignant disease associated with WT1 expression in a patient, comprising the steps of:
 - (a) incubating a first biological sample with one or more of:

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- (i) a WT1 polypeptide;
- a WT1 polynucleotide encoding a WT1 polypeptide; or (ii)
- (iii) antigen-presenting cell that expresses a WT1 polypeptide;

wherein the biological sample comprises CD4+ and/or CD8+ T cells and is obtained from a patient prior to a therapy or immunization, and wherein the incubation is performed under conditions and for a time sufficient to allow specific activation, proliferation and/or lysis of T cells;

- (b) detecting an amount of activation, proliferation and/or lysis of the T cells;
- (c) repeating steps (a) and (b) using a second biological sample comprising CD4+ and/or CD8+ T cells, wherein the second biological sample is obtained from the same patient following therapy or immunization; and
- comparing the amount of activation, proliferation and/or lysis of T cells in the first and second biological samples, and therefrom monitoring the effectiveness of the therapy or immunization in the patient.
- 80. A method according to claim 73 or claim 79, wherein the malignant disease is a cancer or a leukemia.
- 81. A method for inhibiting the development of a malignant disease associated with WT1 expression in a patient, comprising the steps of:
- incubating CD4+ T cells isolated from a patient with one or more (a) of:
 - a WT1 polypeptide; (i)
 - a polynucleotide encoding a WT1 polypeptide; or (ii)
- (iii) an antigen presenting cell that expresses a WT1 polypeptide;

such that the T cells proliferate; and

- (b) administering to the patient an effective amount of the proliferated T cells, and therefrom inhibiting the development of a malignant disease in the patient.
- 82. A method according to claim 81, wherein the malignant disease is a cancer or a leukemia.
- 83. A method according to claim 81, wherein the step of incubating the T cells is repeated one or more times.
- 84. A method for inhibiting the development of a malignant disease associated with WT1 expression in a patient, comprising the steps of:
- (a) incubating CD4⁺ T cells isolated from a patient with one or more of:
 - (i) a WT1 polypeptide;
 - (ii) a polynucleotide encoding a WT1 polypeptide; or
- (iii) an antigen presenting cell that expresses a WT1 polypeptide;

such that the T cells proliferate;

- (b) cloning one or more cells that proliferated in the presence of WT1 polypeptide; and
- (c) administering to the patient an effective amount of the cloned T cells.
- 85. A method according to claim 84, wherein the malignant disease is a cancer or a leukemia.
- 86. A method for inhibiting the development of a malignant disease associated with WT1 expression in a patient, comprising the steps of:

- (a) incubating CD8⁺ T cells isolated from a patient with one or more of:
 - (i) a WT1 polypeptide;
 - (ii) a polynucleotide encoding a WT1 polypeptide; or
 - (iii) an antigen presenting cell expressing a WT1 polypeptide; such that the T cells proliferate; and
- (b) administering to the patient an effective amount of the proliferated T cells, and therefrom inhibiting the development of a malignant disease in the patient.
- 87. A method according to claim 86, wherein the malignant disease is a cancer or a leukemia.
- 88. A method according to claim 86, wherein the step of incubating the T cells is repeated one or more times.
- 89. A method for inhibiting the development of a malignant disease associated with WT1 expression in a patient, comprising the steps of:
- (a) incubating CD8⁺ T cells isolated from a patient with one or more of:
 - (i) a WT1 polypeptide
 - (ii) a polynucleotide encoding a WT1 polypeptide; or
- (iii) an antigen presenting cell that expresses a WT1 polypeptide;

such that the T cells proliferate;

- (b) cloning one or more cells that proliferated in the presence of WT1 polypeptide; and
- (c) administering to the patient an effective amount of the cloned T cells.

- 90. A method according to claim 89, wherein the malignant disease is a cancer or a leukemia.
- 91. A method for determining the presence or absence of a malignant disease associated with WT1 expression in a patient, comprising the steps of:
- (a) incubating CD4⁺ T cells isolated from a patient with one or more of:
 - (i) a WT1 polypeptide;
 - (ii) a polynucleotide encoding a WT1 polypeptide; or
- (iii) an antigen presenting cell that expresses a WT1 polypeptide; and
- (b) detecting the presence or absence of specific activation of the T cells, therefrom determining the presence or absence of a malignant disease associated with WT1 expression.
- 92. A method according to claim 91, wherein the malignant disease is a cancer or a leukemia.
- 93. A method according to claim 91, wherein the step of detecting comprises detecting the presence or absence of proliferation of the T cells.
- 94. A method for determining the presence or absence of a malignant disease associated with WT1 expression in a patient, comprising the steps of:
- (a) incubating CD8⁺ T cells isolated from a patient with a one or more of:
 - (i) a WT1 polypeptide;
 - (ii) a polynucleotide encoding a WT1 polypeptide; or
- (iii) an antigen presenting cell that expresses a WT1 polypeptide; and

- 134
- (b) detecting the presence or absence of specific activation of the T cells, thereby determining the presence or absence of a malignant disease associated with WT1 expression.
- 95. A method according to claim 94, wherein the malignant disease is a cancer or a leukemia.
- 96. A method according to claim 94 wherein the step of detecting comprises detecting the presence or absence of generation of cytolytic activity.
- 97. A method for determining the presence or absence of a malignant disease associated with WT1 expression in a patient, comprising the steps of:
- (a) incubating a biological sample obtained from a patient with one or more of:
 - (i) a WT1 polypeptide;
 - (ii) a polynucleotide encoding a WT1 polypeptide; or
- (iii) an antigen presenting cell that expresses a WT1 polypeptide;

wherein the incubation is performed under conditions and for a time sufficient to allow immunocomplexes to form; and

- (b) detecting immunocomplexes formed between the WT1 polypeptide and antibodies in the biological sample that specifically bind to the WT1 polypeptide; and therefrom determining the presence or absence of a malignant disease associated with WT1 expression.
- 98. A method according to claim 97, wherein the malignant disease is a cancer or a leukemia.
- 99. A method according to claim 97, wherein the step of detecting comprises (a) incubating the immunocomplexes with a detection reagent that is capable

of binding to the immunocomplexes, wherein the detection reagent comprises a reporter group, (b) removing unbound detection reagent, and (c) detecting the presence or absence of the reporter group.

- 100. A method according to claim 99, wherein the detection reagent comprises a second antibody, or antigen-binding fragment thereof, capable of binding to the antibodies that specifically bind to the WT1 polypeptide.
- 101. A method according to claim 99, wherein the detection reagent comprises Protein A.
- 102. A method according to claim 99, wherein the reporter group is selected from the group consisting of radioisotopes, fluorescent groups, luminescent groups, enzymes, biotin and dye particles.
- 103. A method according to claim 97 wherein a reporter group is bound to the WT1 polypeptide, and wherein the step of detecting comprises removing unbound WT1 polypeptide and subsequently detecting the presence or absence of the reporter group.

WO 01/25273 PCT/US00/27465 1/17

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en: Gostes agres die en l'action de la compart de l'action de la compart WC: GCSNEGIGNESDNETN- ITCOMONATELEGALAGIODAEAACAAAAAAAA

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FIG. 2

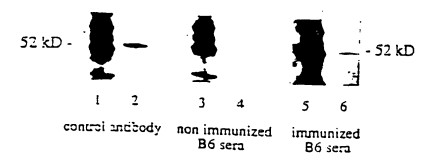


FIG. 3

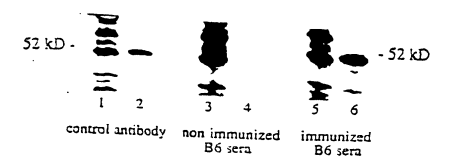
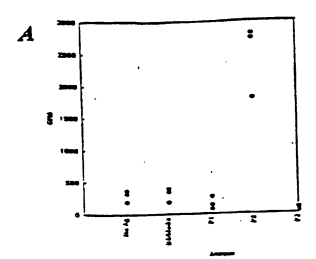
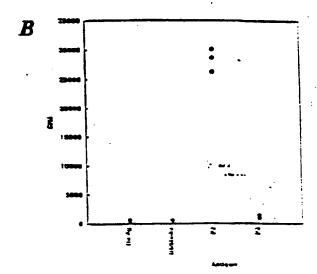


FIG. 4





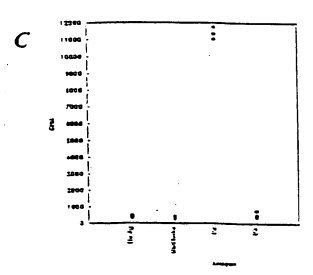


FIG. 5A-5C

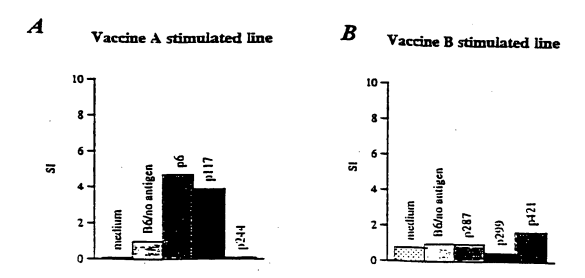


FIG. 6A and 6B

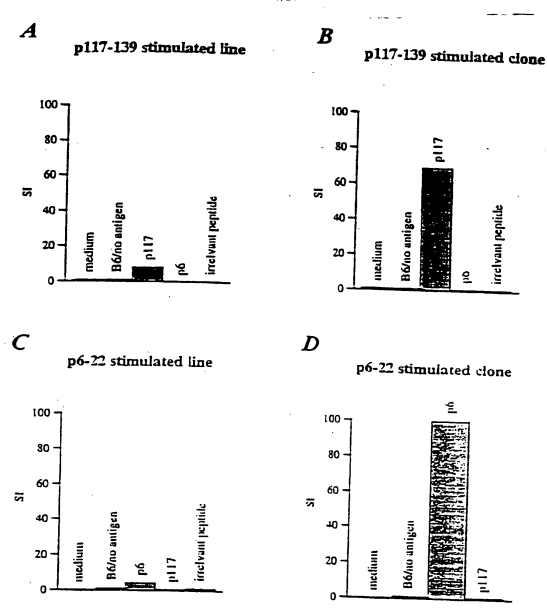
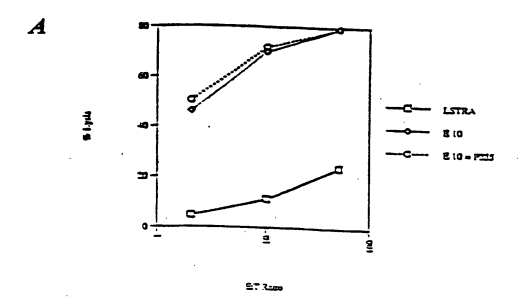


FIG. 7A-7D

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MGSDV															/5
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80	85	5	90	95	100	105	LIG	115	120	125	130	175	140	1 4 6	
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155	160	16	55	170 j	175	130	135	190	195	200	205	210	715	220	200
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FIG. 8A

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80	85	90	95	100	105	110	1:5	120	175	130	176	140	•	
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AHTGVKI														
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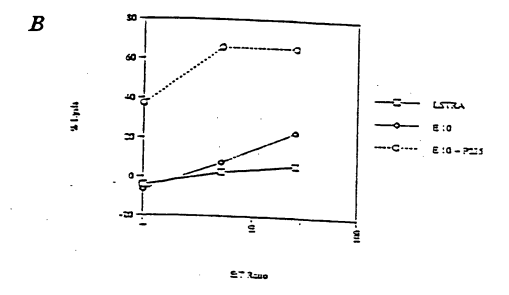


FIG. 9A and 9B

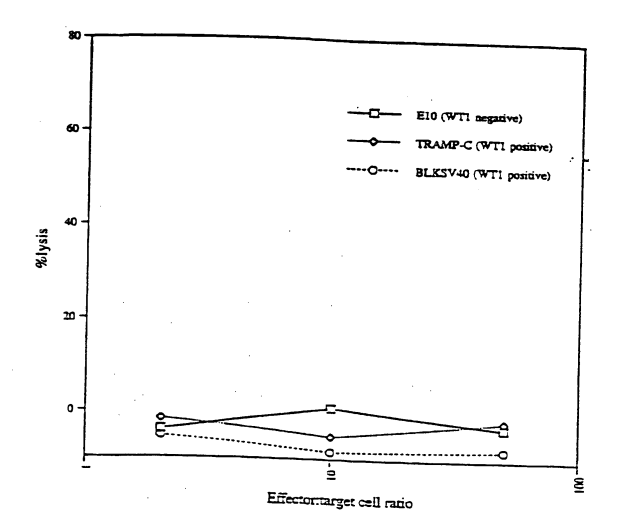


FIG. 10A

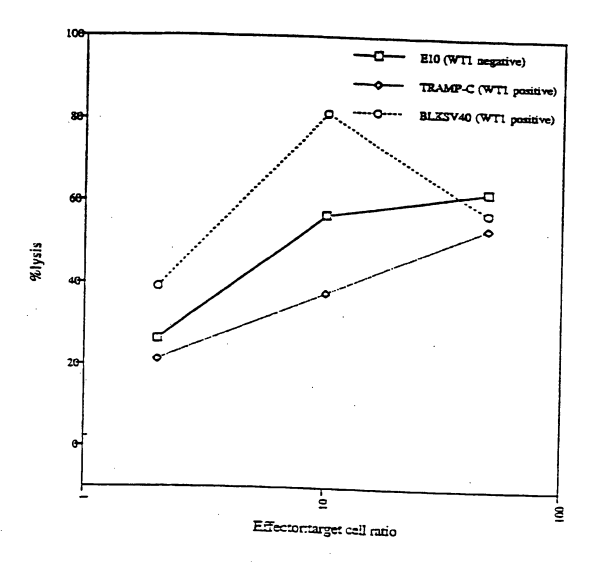


FIG. 10B

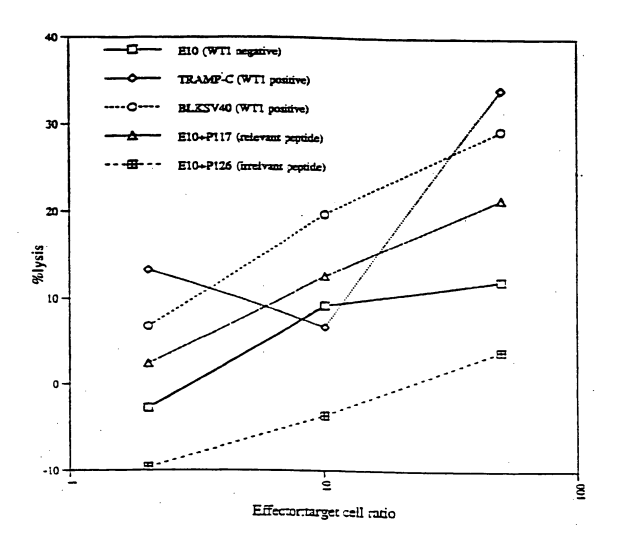


FIG. 10C

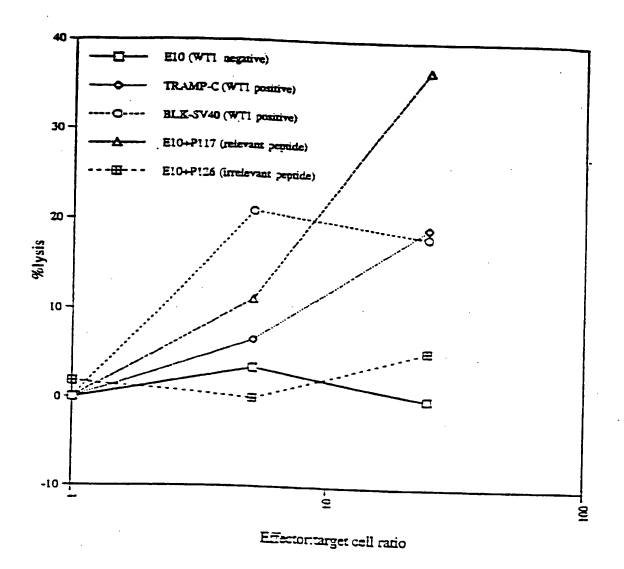
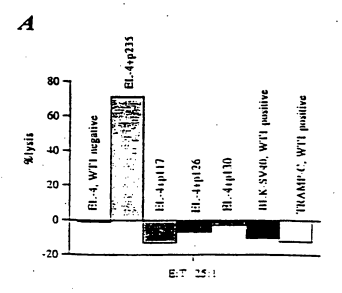


FIG. 10D



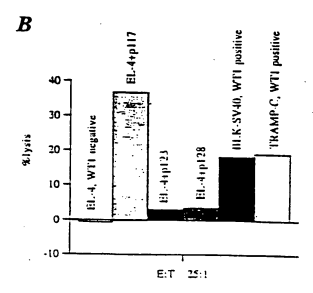
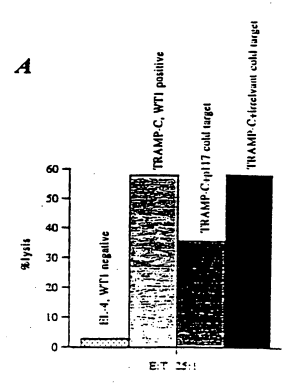


FIG. 11A and 11B



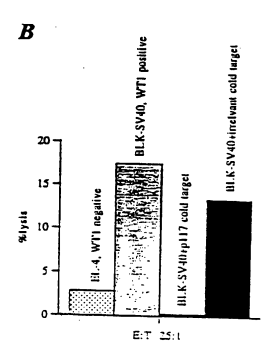
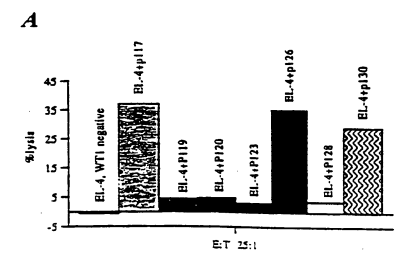


FIG. 12A and 12B



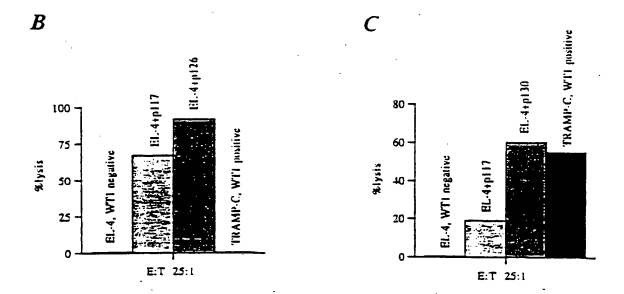


FIG. 13A-13C

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Lys His Ser Gln Pro Trp Gln Val Leu Val Ala Ser Arg Gly Arg Ala
35 40 45

Val Cys Gly Gly Val Leu Val His Pro Gln Trp Val Leu Thr Ala Ala 50 55 60

His Cys Ile Arg Asn Lys Ser Val Ile Leu Leu Gly Arg His Ser Leu 65 70 75 80

Phe His Pro Glu Asp Thr Gly Gln Val Phe Gln Val Ser His Ser Phe 85 90 95

Pro His Pro Leu Tyr Asp Met Ser Leu Leu Lys Asn Arg Phe Leu Arg 100 105 110

Pro Gly Asp Asp Ser Ser His Asp Leu Met Leu Leu Arg Leu Ser Glu 115 120 125

Pro Ala Glu Leu Thr Asp Ala Val Lys Val Met Asp Leu Pro Thr Gln 130 135 140

Glu Pro Ala Leu Gly Thr Thr Cys Tyr Ala Ser Gly Trp Gly Ser Ile 145 150 155 160

Glu Pro Glu Glu Phe Leu Thr Pro Lys Lys Leu Gln Cys Val Asp Leu 165 170 175

His Val Ile Ser Asn Asp Val Cys Ala Gln Val His Pro Gln Lys Val 180 185 190

Thr Lys Phe Met Leu Cys Ala Gly Arg Trp Thr Gly Gly Lys Ser Thr 195 200 205

Cys Ser Gly Asp Ser Gly Gly Pro Leu Val Cys Asn Gly Val Leu Gln 210 215 220

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Ser Leu Tyr Thr Lys Val Val His Tyr Arg Lys Trp Ile Lys Asp Thr 245 250 255

Ile Val Ala Asn Pro 260

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<400> 328

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Lys Glu Leu Lys Phe Val Thr Leu Val Phe Arg His Gly Asp Arg Ser

Pro Ile Asp Thr Phe Pro Thr Asp Pro Ile Lys Glu Ser Ser Trp Pro 50 55 60

Gln Gly Phe Gly Gln Leu Thr Gln Leu Gly Met Glu Gln His Tyr Glu 65 70 75 80

Leu Gly Glu Tyr Ile Arg Lys Arg Tyr Arg Lys Phe Leu Asn Glu Ser 85 90 95

Tyr Lys His Glu Gln Val Tyr Ile Arg Ser Thr Asp Val Asp Arg Thr 100 105 110

Leu Met Ser Ala Met Thr Asn Leu Ala Ala Leu Phe Pro Pro Glu Gly
115 120 125

Val Ser Ile Trp Asn Pro Ile Leu Leu Trp Gln Pro Ile Pro Val His 130 135 140

Thr Val Pro Leu Ser Glu Asp Gln Leu Leu Tyr Leu Pro Phe Arg Asn 145 150 155 160

Cys Pro Arg Phe Gln Glu Leu Glu Ser Glu Thr Leu Lys Ser Glu Glu 165 170 175

Phe Gln Lys Arg Leu His Pro Tyr Lys Asp Phe Ile Ala Thr Leu Gly 180 185 190

Lys Leu Ser Gly Leu His Gly Gln Asp Leu Phe Gly Ile Trp Ser Lys 195 200 205

Val Tyr Asp Pro Leu Tyr Cys Glu Ser Val His Asn Phe Thr Leu Pro 210 215 220

Ser Trp Ala Thr Glu Asp Thr Met Thr Lys Leu Arg Glu Leu Ser Glu 225 230 235 240

Leu Ser Leu Leu Ser Leu Tyr Gly Ile His Lys Gln Lys Glu Lys Ser 245 250 255

Arg Leu Gln Gly Gly Val Leu Val Asn Glu Ile Leu Asn His Met Lys 260 265 270

Arg Ala Thr Gln Ile Pro Ser Tyr Lys Lys Leu Ile Met Tyr Ser Ala 275 280 285

His Asp Thr Thr Val Ser Gly Leu Gln Met Ala Leu Asp Val Tyr Asn

290 295 300 Gly Leu Leu Pro Pro Tyr Ala Ser Cys His Leu Thr Glu Leu Tyr Phe Glu Lys Gly Glu Tyr Phe Val Glu Met Tyr Tyr Arg Asn Glu Thr Gln 330 His Glu Pro Tyr Pro Leu Met Leu Pro Gly Cys Ser Pro Ser Cys Pro Leu Glu Arg Phe Ala Glu Leu Val Gly Pro Val Ile Pro Gln Asp Trp 360 Ser Thr Glu Cys Met Thr Thr Asn Ser His Gln Gly Thr Glu Asp Ser 370 Thr Asp 385 <210> 329 <211> 261 <212> PRT <213> Homo sapiens <400> 329 Met Trp Val Pro Val Val Phe Leu Thr Leu Ser Val Thr Trp Ile Gly 10 Ala Ala Pro Leu Ile Leu Ser Arg Ile Val Gly Gly Trp Glu Cys Glu Lys His Ser Gln Pro Trp Gln Val Leu Val Ala Ser Arg Gly Arg Ala Val Cys Gly Gly Val Leu Val His Pro Gln Trp Val Leu Thr Ala Ala His Cys Ile Arg Asn Lys Ser Val Ile Leu Leu Gly Arg His Ser Leu Phe His Pro Glu Asp Thr Gly Gln Val Phe Gln Val Ser His Ser Phe Pro His Pro Leu Tyr Asp Met Ser Leu Leu Lys Asn Arg Phe Leu Arg 105 Pro Gly Asp Asp Ser Ser His Asp Leu Met Leu Leu Arg Leu Ser Glu 120 Pro Ala Glu Leu Thr Asp Ala Val Lys Val Met Asp Leu Pro Thr Gln 130 Glu Pro Ala Leu Gly Thr Thr Cys Tyr Ala Ser Gly Trp Gly Ser Ile 150

Glu Pro Glu Glu Phe Leu Thr Pro Lys Lys Leu Gln Cys Val Asp Leu

165 170 175

His Val Ile Ser Asn Asp Val Cys Ala Gln Val His Pro Gln Lys Val 180 185 190

Thr Lys Phe Met Leu Cys Ala Gly Arg Trp Thr Gly Gly Lys Ser Thr 195 200 205

Cys Ser Gly Asp Ser Gly Gly Pro Leu Val Cys Asn Gly Val Leu Gln 210 215 220

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Ser Leu Tyr Thr Lys Val Val His Tyr Arg Lys Trp Ile Lys Asp Thr 245 250 255

Ile Val Ala Asn Pro 260

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<212> PRT

<213> Homo sapien

<400> 330

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Phe Thr Glu Trp Ile Glu Lys Thr Val Gln Ala Ser

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200 Thr Cys Val Ala Ala Thr Leu Leu Val Ala Glu Glu Ala Ala Leu Gly 215 220 Pro Thr Glu Pro Ala Glu Gly Leu Ser Ala Pro Ser Leu Ser Pro His 230 235 Cys Cys Pro Cys Arg Ala Arg Leu Ala Phe Arg Asn Leu Gly Ala Leu 245 250 Leu Pro Arg Leu His Gln Leu Cys Cys Arg Met Pro Arg Thr Leu Arg 265 270 Arg Leu Phe Val Ala Glu Leu Cys Ser Trp Met Ala Leu Met Thr Phe 275 280 Thr Leu Phe Tyr Thr Asp Phe Val Gly Glu Gly Leu Tyr Gln Gly Val 295 300 Pro Arg Ala Glu Pro Gly Thr Glu Ala Arg Arg His Tyr Asp Glu Gly 310 315 Val Arg Met Gly Ser Leu Gly Leu Phe Leu Gln Cys Ala Ile Ser Leu 325 330 Val Phe Ser Leu Val Met Asp Arg Leu Val Gln Arg Phe Gly Thr Arg 340 345 Ala Val Tyr Leu Ala Ser Val Ala Ala Phe Pro Val Ala Ala Gly Ala 360 365 Thr Cys Leu Ser His Ser Val Ala Val Val Thr Ala Ser Ala Ala Leu 375 380 Thr Gly Phe Thr Phe Ser Ala Leu Gln Ile Leu Pro Tyr Thr Leu Ala 390 395 Ser Leu Tyr His Arg Glu Lys Gln Val Phe Leu Pro Lys Tyr Arg Gly 405 410 415 Asp Thr Gly Gly Ala Ser Ser Glu Asp Ser Leu Met Thr Ser Phe Leu 420 425 430 Pro Gly Pro Lys Pro Gly Ala Pro Phe Pro Asn Gly His Val Gly Ala 440 Gly Gly Ser Gly Leu Leu Pro Pro Pro Pro Ala Leu Cys Gly Ala Ser 455 460 Ala Cys Asp Val Ser Val Arg Val Val Gly Glu Pro Thr Glu Ala 470 475 Arg Val Val Pro Gly Arg Gly Ile Cys Leu Asp Leu Ala Ile Leu Asp 485 490 Ser Ala Phe Leu Leu Ser Gln Val Ala Pro Ser Leu Phe Met Gly Ser 505 Ile Val Gln Leu Ser Gln Ser Val Thr Ala Tyr Met Val Ser Ala Ala 520 Gly Leu Gly Leu Val Ala Ile Tyr Phe Ala Thr Gln Val Val Phe Asp 535 Lys Ser Asp Leu Ala Lys Tyr Ser Ala 550

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<211> 385

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 catagittci giqciagigg accgi
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ggngcctcac agtatagatc tggtagcaaa gaagaagaaa caaacactga tctctttctg
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ccacccctct gaccctttgg aactcctctg accctttaga acaagcctac ctaatatctg
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attaaacatg gaataaagat ttgtccttaa atataatcta caagaagact ttgatatttg
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                                                                       480
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atcagggacc accetttggg ttgatatttt gettaatetg catettttga gtaagateat
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                                                                        300
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<213> Homo sapiens

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Lys Lys Asp Arg Ala Trp Leu Arg Cys Pro Glu Ala Val Ala Gly Phe

Pro Leu Gly Ser Asp Cys Arg Glu Gly Gly Arg Gln Gly Cys Gly Gly

Ser Asp Asp Glu Asp Asp Leu Gly Val Ala Pro Gly Leu Ala Pro Ala

Trp Ala Leu Thr Gln Pro Pro Ser Gln Ser Pro Gly Pro Gln Ser Leu 105

Pro Ser Thr Pro Ser Ser Ile Trp Pro Gln Trp Val Ile Leu Ile Thr

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Lys His Glu Asp Pro Met Gly Gln Gln Gly Ser Leu Gly Glu Gln Gln 180 185 190

Tyr Ser Val Pro Pro Pro Val Tyr Gly Cys His Thr Pro Thr Asp Ser 195 200 205

Cys Thr Gly Ser Gln Ala Leu Leu Leu Arg Thr Pro Tyr Ser Ser Asp 210 215 220

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Met Asn Leu Gly Ala Thr Leu Lys Gly His Ser Thr Gly Tyr Glu Ser 245 250 255

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